

RESULT 2
US-09-031-626-93
Sequence 93, Application US/09031626
Patent No. 6228581
GENERAL INFORMATION:
APPLICANT: Acton, Susan L.
APPLICANT: O'Donovan, Jose M.
TITLE OF INVENTION: DIAGNOSTIC ASSAYS AND KITS FOR BODY MASS AND
TITLE OF INVENTION: CARDIOVASCULAR DISORDERS
FILE REFERENCE: MIA-005.04
CURRENT APPLICATION NUMBER: US/09/031,626
CURRENT FILING DATE: 1998-02-27
EARLIER APPLICATION NUMBER: 08/890,979
EARLIER FILING DATE: 1997-07-10
NUMBER OF SEQ. ID NOS: 121
SOFTWARE: Patentin Ver. 2.0
SEQ. ID NO 93
LENGTH: 34
TYPE: DNA
ORGANISM: Human

US-09-031-626-93

Query Match 1.2%; Score 31; DB 4; Length 34;
Best Local Similarity 100.0%; Pred. No. 0.0002;
Matches 31; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1085 cctgttctctccatccactctctctcaaa.1115
DB 1 cctgttctctccatccactctctctcaaa 31

RESULT 3

US-09-264-693-7/c
Sequence 7, Application US/09264693
Patent No. 6261760

GENERAL INFORMATION:

APPLICANT: Fielding, Christopher E

APPLICANT: Fielding, Phoebe E

TITLE OF INVENTION: REGULATION OF THE CELL CYCLE BY STEROIDS

FILE REFERENCE: 2500.141US1 Regulation of cell cycle

CURRENT FILING DATE: 1999-03-08

EARLIER APPLICATION NUMBER: 60/077,351

EARLIER FILING DATE: 1998-03-09

NUMBER OF SEQ ID NOS: 10

SOFTWARE: PatentIn Ver. 2.0

SEQ ID NO 7

LENGTH: 30

TYPE: DNA

ORGANISM: Artificial Sequence

FEATURE:

OTHER INFORMATION: Description of Artificial Sequence: Oligo

OTHER INFORMATION: nucleotide probe to CIA-1 mRNA = nucleotides

OTHER INFORMATION: 1514-1543 of human CIA-1 cDNA

US-09-264-693-7

Query Match 1.2%; Score 30; DB 4; Length 30;
Best Local Similarity 100.0%; Pred. No. 0.00055;
Matches 30; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1514 aggataagagagccatcaggctatctg 1543
DB 30 AGGTAAGAGAGCCATCAGGCTATCTCTG 1

RESULT 4

US-08-707-399E-5/c

Sequence 5, Application US/08707399E

Patent No. 6008014

GENERAL INFORMATION:

APPLICANT: Acton, Susan and Gimeno, Carlos

TITLE OF INVENTION: Lipid Metabolic Pathway Compositions

TITLE OF INVENTION: and Therapeutic and Diagnostic Uses Therefor

NUMBER OF SEQUENCES: 23

CORRESPONDENCE ADDRESS:

ADDRESSEE: LAHIVE & COCKFIELD, LLP

STREET: 28 State Street

CITY: Boston

STATE: Massachusetts

COUNTRY: USA

ZIP: 02109

COMPUTER READABLE FORM:

MEDIUM TYPE: Floppy disk

COMPUTER: IBM PC compatible

OPERATING SYSTEM: PC-DOS/MS-DOS

SOFTWARE: PatentIn Release #1.0, Version #1.25

CURRENT APPLICATION DATA:

APPLICATION NUMBER: US/08/707,399E

FILING DATE: September 4, 1996

PRIOR APPLICATION DATA:

APPLICATION NUMBER:

FILING DATE:

ATTORNEY/AGENT INFORMATION:

NAME: Amy E. Mandragouras

REGISTRATION NUMBER: 36,207

REFERENCE/DOCKET NUMBER: MNT-006

TELECOMMUNICATION INFORMATION:

TELEPHONE: (617)227-5941

TELEFAX: (617)227-7400

INFORMATION FOR SEQ ID NO: 5:

SEQUENCE CHARACTERISTICS:

LENGTH: 33 base pairs

TYPE: nucleic acid

STRANDEDNESS: single

TOPOLOGY: linear

MOLECULE TYPE: CDNA

US-08-707-399E-5

Query Match 1.0%; Score 26; DB 3; Length 33;
Best Local Similarity 100.0%; Pred. No. 0.031;
Matches 26; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1576 gtgctgcaggaagcaaacctgtagg 1601
DB 33 GTGCTGCAGGAGCAAAACTGTAGG-8

RESULT 5

US-08-890-980-42/c

Sequence 42, Application US/08890980

Patent No. 5998141

GENERAL INFORMATION:

APPLICANT: Acton, Susan L.

TITLE OF INVENTION: SR-B1 NUCLEIC ACIDS AND USES THEREFOR

NUMBER OF SEQUENCES: 86

CORRESPONDENCE ADDRESS:

ADDRESSEE: FOLEY, HOAG & ELIOT LLP

STREET: One Post Office Square

CITY: Boston

STATE: MA

COUNTRY: USA

ZIP: 02109-2170

COMPUTER READABLE FORM:

MEDIUM TYPE: Floppy disk

COMPUTER: IBM-PC compatible

OPERATING SYSTEM: PC-DOS/MS-DOS

SOFTWARE: PatentIn Release #1.0, Version #1.30

CURRENT APPLICATION DATA:

APPLICATION NUMBER: US/08/890,980

FILING DATE: 10-JUL-1997

CLASSIFICATION: 435

ATTORNEY/AGENT INFORMATION:

NAME: Arnold, Beth E.

REGISTRATION NUMBER: 35,430

REFERENCE/DOCKET NUMBER: MIA-005.01

TELECOMMUNICATION INFORMATION:

TELEPHONE: 617-832-1000

TELEFAX: 617-832-7000

INFORMATION FOR SEQ ID NO: 42:

SEQUENCE CHARACTERISTICS:

LENGTH: 24 base pairs

TYPE: nucleic acid

STRANDEDNESS: single

TOPOLOGY: linear

MOLECULE TYPE: other nucleic acid

DESCRIPTION: /desc = "primer"

Query Match 0.9%; Score 24; DB 2; Length 24;
Best Local Similarity 100.0%; Pred. No. 0.23;
Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

US-08-890-980-42

Db 23 GCTACTGTGGCGCTGTGCTGGCG 1

RESULT 10

US-09-031-626-90/c
Sequence 90, Application US/09031626
Patent No. 6228581
GENERAL INFORMATION:
APPLICANT: Acton, Susan L.
TITLE OF INVENTION: DIAGNOSTIC ASSAYS AND KITS FOR BODY MASS AND
FILE REFERENCE: MIA-005.04
CURRENT APPLICATION NUMBER: US/09/031,626
CURRENT FILING DATE: 1998-02-27
EARLIER APPLICATION NUMBER: 08/890,979
EARLIER FILING DATE: 1997-07-10
NUMBER OF SEQ ID NOS: 121
SOFTWARE: PatentIn Ver. 2.0
SEQ ID NO 90
LENGTH: 23
TYPE: DNA
ORGANISM: Human
US-09-031-626-90

Query Match 0.9%; Score 23; DB 4; Length 23;
Best Local Similarity 100.0%; Pred. No. 0.64;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 123 gctactgtgcgctgtgctggcg 145
Db 23 GCTACTGTGGCGCTGTGCTGGCG 1

RESULT 11

US-08-890-980-72/c
Sequence 72, Application US/08890980
Patent No. 5998141
GENERAL INFORMATION:
APPLICANT: Acton, Susan L.
TITLE OF INVENTION: SR-BI NUCLEIC ACIDS AND USES THEREFOR
NUMBER OF SEQUENCES: 86
CORRESPONDENCE ADDRESS:
ADDRESSEE: FOLEY, HOAG & ELIOT LLP
STREET: One Post Office Square
CITY: Boston
STATE: MA
COUNTRY: USA
ZIP: 02109-2170
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/890,980
FILING DATE: 10-JUL-1997
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Arnold, Beth E.
REGISTRATION NUMBER: 35,430
REFERENCE/DOCKET NUMBER: MIA-005.01
TELECOMMUNICATION INFORMATION:
TELEPHONE: 617-832-1000
TELEFAX: 617-832-7000
INFORMATION FOR SEQ ID NO: 72:
SEQUENCE CHARACTERISTICS:
LENGTH: 31 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear

MOLECULE TYPE: other nucleic acid
DESCRIPTION: /desc = "probe"
US-08-890-980-72

Query Match 0.9%; Score 23; DB 2; Length 31;
Best Local Similarity 100.0%; Pred. No. 0.64;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1112 tcaacgccgaccggtctgcca 1134
Db 23 TCAACGCCGACCGGTTCTGCGCA 1

RESULT 12

US-08-890-980-74
Sequence 74, Application US/08890980
Patent No. 5998141
GENERAL INFORMATION:
APPLICANT: Acton, Susan L.
TITLE OF INVENTION: SR-BI NUCLEIC ACIDS AND USES THEREFOR
NUMBER OF SEQUENCES: 86
CORRESPONDENCE ADDRESS:
ADDRESSEE: FOLEY, HOAG & ELIOT LLP
STREET: One Post Office Square
CITY: Boston
STATE: MA
COUNTRY: USA
ZIP: 02109-2170
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/890,980
FILING DATE: 10-JUL-1997
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Arnold, Beth E.
REGISTRATION NUMBER: 35,430
REFERENCE/DOCKET NUMBER: MIA-005.01
TELECOMMUNICATION INFORMATION:
TELEPHONE: 617-832-1000
TELEFAX: 617-832-7000
INFORMATION FOR SEQ ID NO: 74:
SEQUENCE CHARACTERISTICS:
LENGTH: 31 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: other nucleic acid
DESCRIPTION: /desc = "probe"
US-08-890-980-74

Query Match 0.9%; Score 23; DB 2; Length 31;
Best Local Similarity 100.0%; Pred. No. 0.64;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1112 tcaacgccgaccggtctgcca 1134
Db 9 TCAACGCCGACCGGTTCTGCGCA 31

RESULT 13

US-09-032-894-72/c
Sequence 72, Application US/09032894
Patent No. 6130041
GENERAL INFORMATION:
APPLICANT: Acton, Susan L.
TITLE OF INVENTION: SR-BI NUCLEIC ACIDS AND USES THEREFOR
FILE REFERENCE: MIA-005.03

;; CURRENT APPLICATION NUMBER: US/09/032,894
;; CURRENT FILING DATE: 1998-02-27
;; EARLIER APPLICATION NUMBER: 08/890,980
;; EARLIER FILING DATE: 1997-07-10
;; NUMBER OF SEQ ID NOS: 121
;; SOFTWARE: Patentln Ver. 2.0
;; SEQ ID NO 72
;; LENGTH: 31
;; TYPE: DNA
;; ORGANISM: Human
US-09-032-894-72

Query Match 0.9%; Score 23; DB 3; Length 31;
Best Local Similarity 100.0%; Pred. No. 0.64;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgccgaccggtcttgca 1134
DB 23 TCAACGCCGACCGGTTCTGCA 1

RESULT 14
US-09-032-894-74
;; Sequence 74, Application US/09032894
;; Patent No. 6130041
;; GENERAL INFORMATION:
;; APPLICANT: Acton, Susan L.
;; TITLE OF INVENTION: SR-BI NUCLEIC ACIDS AND USES THEREFOR
;; FILE REFERENCE: MIA-005.03
;; CURRENT APPLICATION NUMBER: US/09/032,894
;; CURRENT FILING DATE: 1998-02-27
;; EARLIER APPLICATION NUMBER: 08/890,980
;; EARLIER FILING DATE: 1997-07-10
;; NUMBER OF SEQ ID NOS: 121
;; SOFTWARE: Patentln Ver. 2.0
;; SEQ ID NO 74
;; LENGTH: 31
;; TYPE: DNA
;; ORGANISM: Human
US-09-032-894-74

Query Match 0.9%; Score 23; DB 3; Length 31;
Best Local Similarity 100.0%; Pred. No. 0.64;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgccgaccggtcttgca 1134
DB 9 tcaacgccgaccggtcttgca 31

RESULT 15
US-09-031-626-72/C
;; Sequence 72, Application US/09031626
;; Patent No. 6228581
;; GENERAL INFORMATION:
;; APPLICANT: Acton, Susan L.
;; APPLICANT: Ordovas, Jose M.
;; TITLE OF INVENTION: DIAGNOSTIC ASSAYS AND KITS FOR BODY MASS AND
;; TITLE OF INVENTION: CARDIOVASCULAR DISORDERS
;; FILE REFERENCE: MIA-005.04
;; CURRENT APPLICATION NUMBER: US/09/031,626
;; CURRENT FILING DATE: 1998-02-27
;; EARLIER APPLICATION NUMBER: 08/890,979
;; EARLIER FILING DATE: 1997-07-10
;; NUMBER OF SEQ ID NOS: 121
;; SOFTWARE: Patentln Ver. 2.0
;; SEQ ID NO 72
;; LENGTH: 31
;; TYPE: DNA
;; ORGANISM: Human
US-09-031-626-72

Query Match 0.9%; Score 23; DB 4; Length 31;
Best Local Similarity 100.0%; Pred. No. 0.64;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgccgaccggtcttgca 1134
DB 23 TCAACGCCGACCGGTTCTGCA 1

RESULT 16
US-09-031-626-74
;; Sequence 74, Application US/09031626
;; Patent No. 6228581
;; GENERAL INFORMATION:
;; APPLICANT: Acton, Susan L.
;; APPLICANT: Ordovas, Jose M.
;; TITLE OF INVENTION: DIAGNOSTIC ASSAYS AND KITS FOR BODY MASS AND
;; TITLE OF INVENTION: CARDIOVASCULAR DISORDERS
;; FILE REFERENCE: MIA-005.04
;; CURRENT APPLICATION NUMBER: US/09/031,626
;; CURRENT FILING DATE: 1998-02-27
;; EARLIER APPLICATION NUMBER: 08/890,979
;; EARLIER FILING DATE: 1997-07-10
;; NUMBER OF SEQ ID NOS: 121
;; SOFTWARE: Patentln Ver. 2.0
;; SEQ ID NO 74
;; LENGTH: 31
;; TYPE: DNA
;; ORGANISM: Human
US-09-031-626-74

Query Match 0.9%; Score 23; DB 4; Length 31;
Best Local Similarity 100.0%; Pred. No. 0.64;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgccgaccggtcttgca 1134
DB 9 tcaacgccgaccggtcttgca 31

RESULT 17
US-08-707-399E-4
;; Sequence 4, Application US/08707399E
;; Patent No. 6008014
;; GENERAL INFORMATION:
;; APPLICANT: Acton, Susan and Gimeno, Carlos
;; TITLE OF INVENTION: Lipid Metabolic Pathway Compositions
;; TITLE OF INVENTION: and Therapeutic and Diagnostic Uses Therefor
;; NUMBER OF SEQUENCES: 23
;; CORRESPONDENCE ADDRESS:
;; ADDRESSEE: LAHIVE & COCKFIELD, LLP
;; STREET: 28 State Street
;; CITY: Boston
;; STATE: Massachusetts
;; COUNTRY: USA
;; ZIP: 02109
;; COMPUTER READABLE FORM:
;; MEDIUM TYPE: Floppy disk
;; COMPUTER: IBM PC compatible
;; OPERATING SYSTEM: PC-DOS/MS-DOS
;; SOFTWARE: Patentln Release #1.0, Version #1.25
;; CURRENT APPLICATION DATA:
;; APPLICATION NUMBER: US/08/707,399E
;; FILING DATE: September 4, 1996
;; PRIOR APPLICATION DATA:
;; APPLICATION NUMBER:
;; FILING DATE:
;; ATTORNEY/AGENT INFORMATION:
;; NAME: Amy E. Mandiragouras
;; REGISTRATION NUMBER: 36,207

REFERENCE/DOCKET NUMBER: MNT-006
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)227-5941
INFORMATION FOR SEQ ID NO: 4:
SEQUENCE CHARACTERISTICS:
LENGTH: 36 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: cdna
US-08-707-399E-4

Query Match 0.7%; Score 18; DB 3; Length 36;
Best Local Similarity 100.0%; Pred. No. 98;
Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1456 caaatccgagccaagag 1473
DB 19 CAAATCCGAGCCCAAGAG 36

RESULT 18
US-09-306-290-12/c
Sequence 12, Application US/09306290
Patent No. 6221635
GENERAL INFORMATION:
APPLICANT: Rovera, Giovanni
APPLICANT: Mukhopadhyay, Sunil
TITLE OF INVENTION: METHODS FOR SOLID-PHASE AMPLIFICATION OF DNA TEMPLATE
TITLE OF INVENTION: (SPAT) USING MULTIBARRAYS
FILE REFERENCE: 09924-10
CURRENT APPLICATION NUMBER: US/09/306,290
CURRENT FILING DATE: 1999-05-06
NUMBER OF SEQ ID NOS: 43
SOFTWARE: PatentIn Ver. 2.0
SEQ ID NO 12
LENGTH: 40
TYPE: DNA
ORGANISM: Artificial Sequence.
FEATURE:
OTHER INFORMATION: Description of Artificial Sequence: Primer RCP
OTHER INFORMATION: RR797
US-09-306-290-12

Query Match 0.7%; Score 18; DB 4; Length 40;
Best Local Similarity 100.0%; Pred. No. 98;
Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2549 atggaagaaaaa 2566
DB 24 ATGGAAGAAAAA 7

RESULT 19
US-08-890-980-71/c
Sequence 71, Application US/08890980
Patent No. 5998141
GENERAL INFORMATION:
APPLICANT: Acton, Susan L.
TITLE OF INVENTION: SR-BI NUCLEIC ACIDS AND USES THEREFOR
NUMBER OF SEQUENCES: 86
CORRESPONDENCE ADDRESS:
ADDRESSEE: FOLEY, HOAG & ELIOT LLP
STREET: One Post Office Square
CITY: Boston.
STATE: MA
COUNTRY: USA
ZIP: 02109-2170
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk

COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/890,980
FILING DATE: 10-JUL-1997
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Arnold, Beth E.
REGISTRATION NUMBER: 35,430
REFERENCE/DOCKET NUMBER: MIA-005.01
TELECOMMUNICATION INFORMATION:
TELEPHONE: 617-832-1000
TELEFAX: 617-832-7000
INFORMATION FOR SEQ ID NO: 71:
SEQUENCE CHARACTERISTICS:
LENGTH: 20 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: other nucleic acid
DESCRIPTION: /desc = "probe"
US-08-890-980-71

Query Match 0.7%; Score 17; DB 12; Length 20;
Best Local Similarity 100.0%; Pred. No. 2,7e+02;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1112 tcaacgccgaccggtt 1128
DB 17 TCAACGCCGACCCGCTT 1

RESULT 20
US-08-890-980-73
Sequence 73, Application US/08890980
Patent No. 5998141
GENERAL INFORMATION:
APPLICANT: Acton, Susan L.
TITLE OF INVENTION: SR-BI NUCLEIC ACIDS AND USES THEREFOR
NUMBER OF SEQUENCES: 86
CORRESPONDENCE ADDRESS:
ADDRESSEE: FOLEY, HOAG & ELIOT LLP
STREET: One Post Office Square
CITY: Boston
STATE: MA
COUNTRY: USA
ZIP: 02109-2170
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/890,980
FILING DATE: 10-JUL-1997
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Arnold, Beth E.
REGISTRATION NUMBER: 35,430
REFERENCE/DOCKET NUMBER: MIA-005.01
TELECOMMUNICATION INFORMATION:
TELEPHONE: 617-832-1000
TELEFAX: 617-832-7000
INFORMATION FOR SEQ ID NO: 73:
SEQUENCE CHARACTERISTICS:
LENGTH: 20 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: other nucleic acid
DESCRIPTION: /desc = "probe"

US-08-890-980-73

Query Match 0.7%; Score 17; DB 2; Length 20;
Best Local Similarity 100.0%; Pred. No. 2.7e+02;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgcgaccgggtt 1128
|||||
DB 4 tcaacgcgaccgggtt 20

RESULT 21

US-09-032-894-71/c
; Sequence 71, Application US/09032894
; Patent No. 6130041
; GENERAL INFORMATION:
; APPLICANT: Acton, Susan L.
; TITLE OF INVENTION: SR-BI NUCLEIC ACIDS AND USES THEREFOR
; FILE REFERENCE: MIA-005.03
; CURRENT APPLICATION NUMBER: US/09/032.894
; EARLIER FILING DATE: 1998-02-27
; EARLIER APPLICATION NUMBER: 08/890.980
; EARLIER FILING DATE: 1997-07-10
; NUMBER OF SEQ ID NOS: 121
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 71
; LENGTH: 20
; TYPE: DNA
; ORGANISM: Human
US-09-032-894-71

Query Match 0.7%; Score 17; DB 3; Length 20;
Best Local Similarity 100.0%; Pred. No. 2.7e+02;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgcgaccgggtt 1128
|||||
DB 17 tcaacgcgaccgggtt 1

RESULT 22

US-09-032-894-73
; Sequence 73, Application US/09032894
; Patent No. 6130041
; GENERAL INFORMATION:
; APPLICANT: Acton, Susan L.
; TITLE OF INVENTION: SR-BI NUCLEIC ACIDS AND USES THEREFOR
; FILE REFERENCE: MIA-005.03
; CURRENT APPLICATION NUMBER: US/09/032.894
; EARLIER FILING DATE: 1998-02-27
; EARLIER APPLICATION NUMBER: 08/890.980
; EARLIER FILING DATE: 1997-07-10
; NUMBER OF SEQ ID NOS: 121
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 73
; LENGTH: 20
; TYPE: DNA
; ORGANISM: Human
US-09-032-894-73

Query Match 0.7%; Score 17; DB 3; Length 20;
Best Local Similarity 100.0%; Pred. No. 2.7e+02;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgcgaccgggtt 1128
|||||
DB 4 tcaacgcgaccgggtt 20

RESULT 23

US-09-031-626-71/c
; Sequence 71, Application US/09031626
; Patent No. 6228581
; GENERAL INFORMATION:
; APPLICANT: Acton, Susan L.
; APPLICANT: Ordovas, Jose M.
; TITLE OF INVENTION: DIAGNOSTIC ASSAYS AND KITS FOR BODY MASS AND
; FILE REFERENCE: MIA-005.04
; CURRENT APPLICATION NUMBER: US/09/031.626
; EARLIER FILING DATE: 1998-02-27
; EARLIER APPLICATION NUMBER: 08/890.979
; EARLIER FILING DATE: 1997-07-10
; NUMBER OF SEQ ID NOS: 121
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 71
; LENGTH: 20
; TYPE: DNA
; ORGANISM: Human
US-09-031-626-71

Query Match 0.7%; Score 17; DB 4; Length 20;
Best Local Similarity 100.0%; Pred. No. 2.7e+02;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgcgaccgggtt 1128
|||||
DB 17 tcaacgcgaccgggtt 1

RESULT 24

US-09-031-626-73
; Sequence 73, Application US/09031626
; Patent No. 6228581
; GENERAL INFORMATION:
; APPLICANT: Acton, Susan L.
; APPLICANT: Ordovas, Jose M.
; TITLE OF INVENTION: DIAGNOSTIC ASSAYS AND KITS FOR BODY MASS AND
; FILE REFERENCE: MIA-005.04
; CURRENT APPLICATION NUMBER: US/09/031.626
; EARLIER FILING DATE: 1998-02-27
; EARLIER APPLICATION NUMBER: 08/890.979
; EARLIER FILING DATE: 1997-07-10
; NUMBER OF SEQ ID NOS: 121
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 73
; LENGTH: 20
; TYPE: DNA
; ORGANISM: Human
US-09-031-626-73

Query Match 0.7%; Score 17; DB 4; Length 20;
Best Local Similarity 100.0%; Pred. No. 2.7e+02;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgcgaccgggtt 1128
|||||
DB 4 tcaacgcgaccgggtt 20

RESULT 25

US-08-928-465-1/c
; Sequence 1, Application US/08928465
; Patent No. 6204024
; GENERAL INFORMATION:
; APPLICANT: Romano, Joseph
; APPLICANT: Lee, Eun Mi
; TITLE OF INVENTION: CCR5 RNA Transcription Based
; TITLE OF INVENTION: Amplification Assay
; NUMBER OF SEQUENCES: 10

;; CORRESPONDENCE ADDRESS:
;; ADDRESSEE: Akzo No. 6204024e1 Patent Department
;; STREET: 1300 Piccard Drive
;; CITY: Rockville
;; STATE: Maryland
;; COUNTRY: US
;; ZIP: 20850
;;
;; COMPUTER READABLE FORM:
;; MEDIUM TYPE: Floppy disk
;; COMPUTER: IBM PC compatible
;; OPERATING SYSTEM: PC-DOS/MS-DOS
;; SOFTWARE: Patentin Release #1.0, Version #1.30
;; CURRENT APPLICATION DATA:
;; APPLICATION NUMBER: US/08/928,465
;; FILING DATE:
;; CLASSIFICATION: 435
;; ATTORNEY/AGENT INFORMATION:
;; NAME: Gormley, Mary E.
;; REGISTRATION NUMBER: 34,409
;; TELECOMMUNICATION INFORMATION:
;; TELEPHONE: 301-948-7400
;; TELEFAX: 301-948-9751
;; INFORMATION FOR SEQ ID NO: 1:
;; SEQUENCE CHARACTERISTICS:
;; LENGTH: 47 base pairs
;; TYPE: nucleic acid
;; STRANDEDNESS: single
;; TOPOLOGY: not relevant
;; MOLECULE TYPE: other nucleic acid
;; DESCRIPTION: /desc = "DNA oligonucleotide"
;; HYPOTHETICAL: NO
;; FEATURE:
;; NAME/KEY: misc.feature
;; LOCATION: 1..25
;; OTHER INFORMATION: /label= T7 RNA Polymere
;;
US-08-928-465-1
;;
Query Match 0.7%; Score 17; DB 4; Length 47;
Best Local Similarity 100.0%; Pred. No. 2.7e+02;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
OY 1295 tggctcctgcgcgtcctc 1311
|||||
DB 41 TGGTCTGCGCGTCTC 25
;;
RESULT 26
US-09-437-076-1
;; Sequence 1, Application US/09437076
;; Patent No. 6261779
;; GENERAL INFORMATION:
;; APPLICANT: Barber-Guilllem, Emilio
;; APPLICANT: Nelson, M. Bud
;; TITLE OF INVENTION: Nanocrystals having polynucleotide strands and their use to form
;; CURRENT APPLICATION NUMBER: US/09/437,076
;; EARLIER FILING DATE:
;; NUMBER OF SEQ ID NOS: 6
;; SOFTWARE: word for windows
;; SEQ ID NO 1
;; LENGTH: 18
;; TYPE: DNA
;; ORGANISM: Artificial sequence
;; FEATURE:
;; NAME/KEY:
;; LOCATION:
;; OTHER INFORMATION: synthesized
US-09-437-076-1

Query Match 0.6%; Score 16; DB 4; Length 18;
Best Local Similarity 100.0%; Pred. No. 7.5e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
OY 2551 ggaataaaaaaaaaa 2566
|||||
DB 2 ggaataaaaaaaaaa 17
;;
RESULT 27
US-08-928-465-4/C
;; Sequence 4, Application US/08928465
;; Patent No. 6204024
;; GENERAL INFORMATION:
;; APPLICANT: Romano, Joseph
;; APPLICANT: Lee, Eun M.
;; TITLE OF INVENTION: CCR5 RNA Transcription Based
;; NUMBER OF SEQUENCES: 10
;; CORRESPONDENCE ADDRESS:
;; ADDRESSEE: Akzo No. 6204024e1 Patent Department
;; STREET: 1300 Piccard Drive
;; CITY: Rockville
;; STATE: Maryland
;; COUNTRY: US
;; ZIP: 20850
;;
;; COMPUTER READABLE FORM:
;; MEDIUM TYPE: Floppy disk
;; COMPUTER: IBM PC compatible
;; OPERATING SYSTEM: PC-DOS/MS-DOS
;; SOFTWARE: Patentin Release #1.0, Version #1.30
;; CURRENT APPLICATION DATA:
;; APPLICATION NUMBER: US/08/928,465
;; FILING DATE:
;; CLASSIFICATION: 435
;; ATTORNEY/AGENT INFORMATION:
;; NAME: Gormley, Mary E.
;; REGISTRATION NUMBER: 34,409
;; TELECOMMUNICATION INFORMATION:
;; TELEPHONE: 301-948-7400
;; TELEFAX: 301-948-9751
;; INFORMATION FOR SEQ ID NO: 4:
;; SEQUENCE CHARACTERISTICS:
;; LENGTH: 22 base pairs
;; TYPE: nucleic acid
;; STRANDEDNESS: single
;; TOPOLOGY: not relevant
;; MOLECULE TYPE: other nucleic acid
;; DESCRIPTION: /desc = "DNA oligonucleotide"
;; HYPOTHETICAL: NO
;;
US-08-928-465-4
;;
Query Match 0.6%; Score 16; DB 4; Length 22;
Best Local Similarity 100.0%; Pred. No. 7.5e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
OY 1295 tggctcctgcgcgtcctc 1310
|||||
DB 16 TGGTCTGCGCGTCTC 1
;;
RESULT 28
US-08-991-347-5
;; Sequence 5, Application US/08991347
;; Patent No. 6107032
;; GENERAL INFORMATION:
;; APPLICANT: PBO, Svante
;; APPLICANT: Kilger, Christian
;; TITLE OF INVENTION: METHOD FOR THE DIRECT EXPONENTIAL AMPLIFICATION AND
;; TITLE OF INVENTION: SEQUENCING OF DNA MOLECULES AND ITS APPLICATION
;; FILE REFERENCE: 1614-7089
;; CURRENT APPLICATION NUMBER: US/08/991,347

;; CURRENT FILING DATE: 1997-12-16
;; EARLIER APPLICATION NUMBER: DE 19653439.9
;; EARLIER FILING DATE: 1996-12-20
;; NUMBER OF SEQ ID NOS: 6
;; SOFTWARE: Patentin Ver. 2.0
;; SEQ ID NO 5
;; LENGTH: 25
;; TYPE: DNA
;; ORGANISM: Artificial Sequence
;; FEATURE:
;; OTHER INFORMATION: Description of Artificial Sequence:PRIMER
US-08-991-184-5

Query Match 0.6%; Score 16; DB 3; Length 25;
Best Local Similarity 100.0%; Pred No. 7.5e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1295 tggctctgcgcgtcgt 1310
|||||
DB 4 tggctctgcgcgtcgt 19

RESULT 29
US-08-991-184-1
;; Sequence 1, Application US/08991184
;; Patent No. 6225092
;; GENERAL INFORMATION:

;; APPLICANT: P BO, Svante
;; TITLE OF INVENTION: Method for uncoupled, direct, exponential
;; TITLE OF INVENTION: amplification and sequencing of DNA molecules with the additio
;; TITLE OF INVENTION: thermostable DNA polymerase and its application
;; NUMBER OF SEQUENCES: 5
;; CORRESPONDENCE ADDRESS:
;; ADDRESSEE: Nixaido, Marmelstein, Murray & Oram LLP
;; STREET: 655 Fifteenth Street, N.W.; Suite 330
;; CITY: Washington
;; STATE: D.C.
;; COUNTRY: U.S.A.
;; ZIP: 20005-5701

;; COMPUTER READABLE FORM:
;; MEDIUM TYPE: IBM PC compatible
;; OPERATING SYSTEM: PC-DOS/MS-DOS
;; SOFTWARE: Patentin Release #1.0, Version #1.30
;; CURRENT APPLICATION DATA:
;; APPLICATION NUMBER: US/08/991,184
;; FILING DATE: 16-DEC-1997
;; CLASSIFICATION: 435
;; PRIOR APPLICATION DATA:
;; APPLICATION NUMBER: DE 196 53 494.1
;; FILING DATE: 20-DEC-1996
;; CLASSIFICATION: 435
;; ATTORNEY/AGENT INFORMATION:
;; NAME: Wong, King L.
;; REGISTRATION NUMBER: 37,500
;; REFERENCE/DOCKET NUMBER: 1614-7090
;; TELECOMMUNICATION INFORMATION:
;; TELEPHONE: (202) 638-5000
;; TELEFAX: (202) 638-4810
;; INFORMATION FOR SEQ ID NO: 1:
;; SEQUENCE CHARACTERISTICS:
;; LENGTH: 25 base pairs
;; TYPE: nucleic acid
;; STRANDEDNESS: single
;; TOPOLOGY: linear
;; MOLECULE TYPE: other nucleic acid
;; DESCRIPTION: /desc = "oligonucleotide"
US-08-991-184-1

Query Match 0.6%; Score 16; DB 4; Length 25;

Best Local Similarity 100.0%; Pred. No. 7.5e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1295 tggctctgcgcgtcgt 1310
|||||
DB 4 TGGCTCTGCCGCTGCT 19

RESULT 30
US-08-602-716A-1/c
;; Sequence 1, Application US/08602716A
;; Patent No. 596264
;; GENERAL INFORMATION:

;; APPLICANT: FRIEDHOFF, Arnold J.
;; APPLICANT: BASHAM, Daryl A.
;; APPLICANT: MILLER, Jeanette C.
;; TITLE OF INVENTION: PSYCHOSIS PROTECTING NUCLEIC ACID,
;; TITLE OF INVENTION: PEPTIDES, COMPOSITIONS AND METHODS OF USE
;; NUMBER OF SEQUENCES: 12
;; CORRESPONDENCE ADDRESS:
;; ADDRESSEE: BROWDY AND NEWMARK
;; STREET: 419 Seventh Street, N.W., Suite 300
;; CITY: Washington
;; STATE: D.C.
;; COUNTRY: USA
;; ZIP: 20004

;; COMPUTER READABLE FORM:
;; MEDIUM TYPE: floppy disk
;; OPERATING SYSTEM: PC-DOS/MS-DOS
;; SOFTWARE: Patentin Release #1.0, Version #1.30
;; CURRENT APPLICATION DATA:
;; APPLICATION NUMBER: US/08/602,716A
;; FILING DATE: 23-FEB-1996
;; CLASSIFICATION: 435
;; PRIOR APPLICATION DATA:
;; APPLICATION NUMBER: US 08/060,560
;; FILING DATE: 13-MAY-1993
;; PRIOR APPLICATION DATA: PCT/US94/05545
;; APPLICATION NUMBER: PCT/US94/05545
;; FILING DATE: 13-MAY-1994
;; ATTORNEY/AGENT INFORMATION:
;; NAME: BROWDY, Roger L.
;; REGISTRATION NUMBER: 25,618
;; REFERENCE/DOCKET NUMBER: FRIEDHOFF-1A
;; TELECOMMUNICATION INFORMATION:
;; TELEPHONE: 202-628-5197
;; TELEFAX: 202-737-3528
;; INFORMATION FOR SEQ ID NO: 1:
;; SEQUENCE CHARACTERISTICS:
;; LENGTH: 26 base pairs
;; TYPE: nucleic acid
;; STRANDEDNESS: single
;; TOPOLOGY: linear
;; MOLECULE TYPE: CDNA
US-08-602-716A-1

Query Match 0.6%; Score 16; DB 2; Length 26;
Best Local Similarity 100.0%; Pred. No. 7.5e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaaaa 2566
|||||
DB 18 GGAATAAAAAAAAAA 3

RESULT 31
US-07-862-831A-15
;; Sequence 15, Application US/07862831A
;; Patent No. 5356802
;; GENERAL INFORMATION:
;; APPLICANT: Chandrasegaran, Srinivasan

TITLE OF INVENTION: Functional Domains in Foki Restriction
TITLE OF INVENTION: Endonuclease
NUMBER OF SEQUENCES: 20
CORRESPONDENCE ADDRESSES:
ADDRESSEE: Cushman, Darby & Cushman
STREET: 1615 L St., N.W.
CITY: Washington
STATE: D.C.
COUNTRY: USA
ZIP: 20036-5601
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/07/862,831A
FILING DATE: 19920403
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Kokulis, Paul N.
REGISTRATION NUMBER: 16,773
REFERENCE/DOCKET NUMBER: PNK/4130/93738/SLO
TELECOMMUNICATION INFORMATION:
TELEPHONE: 202-861-3000
TELEFAX: 202-822-0944
TELEX: 6714627 CUSH
INFORMATION FOR SEQ ID NO: 15:
SEQUENCE CHARACTERISTICS:
LENGTH: 30 base pairs
TYPE: NUCLEIC ACID
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: DNA (genomic)
US-07-862-831A-15

Query Match 0.6%; Score:16; DB 1; Length 30;
Best Local Similarity 100.0%; Pred.No.7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaataaa 2566
|||||

Db 14 GGAATAAATAAATAA 29

RESULT 32
US-08-126-564A-15
Sequence 15, Application US/08126564A
Patent No. 5436150
GENERAL INFORMATION:
APPLICANT: Chandrasegaran, Srinivasan
TITLE OF INVENTION: Functional Domains in Foki
TITLE OF INVENTION: Restriction Endonuclease
NUMBER OF SEQUENCES: 48
CORRESPONDENCE ADDRESSES:
ADDRESSEE: Cushman, Darby & Cushman
STREET: 1100 New York Ave., N.W.
CITY: Washington
STATE: D.C.
COUNTRY: USA
ZIP: 20005-3918
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0,
SOFTWARE: Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/126,564A
FILING DATE: 27-SEPTEMBER-93
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:

NAME: Kokulis, Paul N.
REGISTRATION NUMBER: 16,773
REFERENCE/DOCKET NUMBER: PNK/4130/82506/CLB
TELECOMMUNICATION INFORMATION:
TELEPHONE: 202-861-3503
TELEFAX: 202-822-0944
TELEX: 6714627 CUSH
INFORMATION FOR SEQ ID NO: 15:
SEQUENCE CHARACTERISTICS:
LENGTH: 30 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: DNA (genomic)
US-08-126-564A-15

Query Match 0.6%; Score 16; DB 1; Length 30;
Best Local Similarity 100.0%; Pred.No.7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaataaa 2566
|||||

Db 14 GGAATAAATAAATAA 29

RESULT 33
PCT-US94-09143-15
Sequence 15, Application PCT/US9409143
GENERAL INFORMATION:
APPLICANT: Chandrasegaran, Srinivasan
TITLE OF INVENTION: Functional Domains in Foki
TITLE OF INVENTION: Restriction Endonuclease
NUMBER OF SEQUENCES: 48
CORRESPONDENCE ADDRESSES:
ADDRESSEE: Cushman, Darby & Cushman
STREET: 1100 New York Ave., N.W.
CITY: Washington
STATE: D.C.
COUNTRY: USA
ZIP: 20005-3918
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0,
SOFTWARE: Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: PCT/US94/09143
FILING DATE: 23-AUG-1994
CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/126,564
FILING DATE: 27-SEPTEMBER-93
ATTORNEY/AGENT INFORMATION:
NAME: Kokulis, Paul N.
REGISTRATION NUMBER: 16,773
REFERENCE/DOCKET NUMBER: PNK/4130/82506/CLB
TELECOMMUNICATION INFORMATION:
TELEPHONE: 202-861-3503
TELEFAX: 202-822-0944
TELEX: 6714627 CUSH
INFORMATION FOR SEQ ID NO: 15:
SEQUENCE CHARACTERISTICS:
LENGTH: 30 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: DNA (genomic)
PCT-US94-09143-15

Query Match 0.6%; Score 16; DB 5; Length 30;

Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaaaaa 2566
|||||
DB 14 GGAATAAAAAAAAA 29

RESULT 34

US-08-324-243-4

Sequence 4, Application US/08324243

Patent No. 5786464

GENERAL INFORMATION:

APPLICANT: SEED, BRIAN

TITLE OF INVENTION: OVEREXPRESSION OF MAMMALIAN AND VIRAL

TITLE OF INVENTION: PROTEINS

NUMBER OF SEQUENCES: 37

CORRESPONDENCE ADDRESS:

ADDRESSEE: Fish & Richardson

STREET: 225 Franklin Street

CITY: Boston

STATE: Massachusetts

COUNTRY: U.S.A.

ZIP: 02110-2804

COMPUTER READABLE FORM:

MEDIUM TYPE: Floppy disk

COMPUTER: IBM PC compatible

OPERATING SYSTEM: PC-DOS/MS-DOS

SOFTWARE: Patentin Release #1.0, Version #1.30B

CURRENT APPLICATION DATA:

APPLICATION NUMBER: US/08/324,243

FILING DATE: 19-SEP-1994

ATTORNEY/AGENT INFORMATION:

NAME: CLARK, PAUL T

REGISTRATION NUMBER: 30,162

REFERENCE/DOCKET NUMBER: 00786/226001

TELECOMMUNICATION INFORMATION:

TELEPHONE: (617) 542-5070

TELEFAX: (617) 542-8906

TELEX: 200154

INFORMATION FOR SEQ ID NO: 4:

SEQUENCE CHARACTERISTICS:

LENGTH: 33 base pairs

TYPE: nucleic acid

STRANDEDNESS: single

TOPOLOGY: linear

US-08-324-243-4

Query Match 0.6%; Score 16; DB 1; Length 33;
Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 225 cttcaacatgtggaag 240
|||||
DB 10 CTTCAACATGTGGAAG 25

RESULT 35
US-08-532-390-4
Sequence 4, Application US/08532390
Patent No. 5795737
GENERAL INFORMATION:
APPLICANT: SEED, BRIAN
APPLICANT: HAAS, JURGEN
TITLE OF INVENTION: High Level Expression of Proteins
NUMBER OF SEQUENCES: 40
CORRESPONDENCE ADDRESS:
ADDRESSEE: Fish & Richardson P.C.
STREET: 225 Franklin Street
CITY: Boston
STATE: Massachusetts
COUNTRY: U.S.A.

ZIP: 02110-2804
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.30B
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/532,390
FILING DATE: 22-SEP-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/324,243
FILING DATE: 19-SEP-1994
ATTORNEY/AGENT INFORMATION:
NAME: LECH, KAREN F.
REGISTRATION NUMBER: 35,238
REFERENCE/DOCKET NUMBER: 00786/294001
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617) 542-5070
TELEFAX: (617) 542-8906
TELEX: 200154
INFORMATION FOR SEQ ID NO: 4:
SEQUENCE CHARACTERISTICS:
LENGTH: 33 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
US-08-532-390-4

Query Match 0.6%; Score 16; DB 1; Length 33;
Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 225 cttcaacatgtggaag 240
|||||
DB 10 CTTCAACATGTGGAAG 25

RESULT 36
US-08-717-294-4
Sequence 4, Application US/08717294
Patent No. 6114148
GENERAL INFORMATION:
APPLICANT: SEED, BRIAN
APPLICANT: HAAS, JURGEN
TITLE OF INVENTION: HIGH LEVEL EXPRESSION OF
NUMBER OF SEQUENCES: 110
CORRESPONDENCE ADDRESS:
ADDRESSEE: Clark & Elbing LLP
STREET: 176 Federal Street
CITY: Boston
STATE: MA
COUNTRY: USA
ZIP: 02110
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FastSeq for Windows Version 2.0
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/717,294
FILING DATE: 20-SEP-1996
CLASSIFICATION: 435
PRIOR APPLICATION NUMBER:
APPLICATION NUMBER:
FILING DATE:
ATTORNEY/AGENT INFORMATION:
NAME: Elbing, Karen L.
REGISTRATION NUMBER: 35,238
REFERENCE/DOCKET NUMBER: 00786/345001
TELECOMMUNICATION INFORMATION:
TELEPHONE: 617-428-0200

TELEFAX: 617-428-7045
TELEX: 4:
INFORMATION FOR SEQ ID NO: 4:
SEQUENCE CHARACTERISTICS:
LENGTH: 33 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: Other
US-08-717-294-4

Query Match 0.6%; Score 16; DB 3; Length 33;
Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 225 ctcaacatgtggaag 240
|||||
DB 10 CTCACACATGTGGAAG 25

RESULT 37
PCT-US95-11511-4
Sequence 4, Application PC/TUS9511511
GENERAL INFORMATION:
APPLICANT: SEED, BRIAN
TITLE OF INVENTION: OVEREXPRESSION OF MAMMALIAN AND VIRAL
PROTEINS
NUMBER OF SEQUENCES: 37
CORRESPONDENCE ADDRESSES:
ADDRESSEE: Fish & Richardson
STREET: 225 Franklin Street
CITY: Boston
STATE: Massachusetts
COUNTRY: U.S.A.
ZIP: 02110-2804
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30B
CURRENT APPLICATION DATA:
APPLICATION NUMBER: PCT/US95/11511
FILING DATE:
ATTORNEY/AGENT INFORMATION:
NAME: CLARK, PAUL T
REGISTRATION NUMBER: 30,162
REFERENCE/DOCKET NUMBER: 00786/226001
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617) 542-5070
TELEFAX: (617) 542-8906
TELEX: 200154
INFORMATION FOR SEQ ID NO: 4:
SEQUENCE CHARACTERISTICS:
LENGTH: 33 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
PCT-US95-11511-4

Query Match 0.6%; Score 16; DB 5; Length 33;
Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 225 ctcaacatgtggaag 240
|||||
DB 10 CTCACACATGTGGAAG 25

RESULT 38
US-08-324-243-3/c
Sequence 3, Application US/08324243

Patent No. 5786464
GENERAL INFORMATION:
APPLICANT: SEED, BRIAN
TITLE OF INVENTION: OVEREXPRESSION OF MAMMALIAN AND VIRAL
PROTEINS
NUMBER OF SEQUENCES: 37
CORRESPONDENCE ADDRESSES:
ADDRESSEE: Fish & Richardson
STREET: 225 Franklin Street
CITY: Boston
STATE: Massachusetts
COUNTRY: U.S.A.
ZIP: 02110-2804
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30B
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/324,243
FILING DATE: 19-SEP-1994
ATTORNEY/AGENT INFORMATION:
NAME: CLARK, PAUL T
REGISTRATION NUMBER: 30,162
REFERENCE/DOCKET NUMBER: 00786/226001
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617) 542-5070
TELEFAX: (617) 542-8906
TELEX: 200154
INFORMATION FOR SEQ ID NO: 3:
SEQUENCE CHARACTERISTICS:
LENGTH: 34 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
US-08-324-243-3

Query Match 0.6%; Score 16; DB 1; Length 34;
Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 225 ctcaacatgtggaag 240
|||||
DB 29 CTCACACATGTGGAAG 14

RESULT 39
US-08-532-390-3/c
Sequence 3, Application US/08532390
Patent No. 5795737
GENERAL INFORMATION:
APPLICANT: SEED, BRIAN
HAAS, JURGEN
TITLE OF INVENTION: High Level Expression of Proteins
NUMBER OF SEQUENCES: 40
CORRESPONDENCE ADDRESSES:
ADDRESSEE: Fish & Richardson P.C.
STREET: 225 Franklin Street
CITY: Boston
STATE: Massachusetts
COUNTRY: U.S.A.
ZIP: 02110-2804
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30B
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/532,390
FILING DATE: 22-SEP-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/324,243

FILING DATE: 19-SEP-1994
ATTORNEY/AGENT INFORMATION:
NAME: LECH, KAREN F.
REGISTRATION NUMBER: 35,238
REFERENCE/DOCKET NUMBER: 00786/294001
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617) 542-5070
TELEFAX: (617) 542-8906
TELEX: 200154
INFORMATION FOR SEQ ID NO: 3:
SEQUENCE CHARACTERISTICS:
LENGTH: 34 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
US-08-532-390-3

Query Match 0.6%; Score 16; DB 1; Length 34;
Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 225 cttcaacatgtggaag 240
DB 29 CTTCAACATGTGGAAG 14

RESULT 40
US-08-717-294-3/c
Sequence 3, Application US/08717294
Patent No. 6114148
GENERAL INFORMATION:
APPLICANT: SEED, BRIAN
TITLE OF INVENTION: HIGH LEVEL EXPRESSION OF
NUMBER OF SEQUENCES: 110
CORRESPONDENCE ADDRESS:
ADDRESSEE: Clark & Elbing LLP
STREET: 176 Federal Street
CITY: Boston
STATE: MA
COUNTRY: USA
ZIP: 02110
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
OPERATING SYSTEM: DOS
SOFTWARE: FASTSEQ for Windows Version 2.0
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/717,294
FILING DATE: 20-SEP-1996
CLASSIFICATION: 435
PRIOR APPLICATION NUMBER:
FILING DATE:
ATTORNEY/AGENT INFORMATION:
NAME: Elbing, Karen L.
REGISTRATION NUMBER: 35,238
REFERENCE/DOCKET NUMBER: 00786/345001
TELECOMMUNICATION INFORMATION:
TELEPHONE: 617-428-0200
TELEFAX: 617-428-7045
TELEX:
INFORMATION FOR SEQ ID NO: 3:
SEQUENCE CHARACTERISTICS:
LENGTH: 34 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: Other
US-08-717-294-3

Query Match 0.6%; Score 16; DB 3; Length 34;
Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 225 cttcaacatgtggaag 240
DB 29 CTTCAACATGTGGAAG 14

RESULT 41
PCT-US95-11511-3/c
Sequence 3, Application PC/TUS9511511
GENERAL INFORMATION:
APPLICANT: SEED, BRIAN
TITLE OF INVENTION: OVEREXPRESSION OF MAMMALIAN AND VIRAL
NUMBER OF SEQUENCES: 37
CORRESPONDENCE ADDRESS:
ADDRESSEE: Fish & Richardson
STREET: 225 Franklin Street
CITY: Boston
STATE: Massachusetts
COUNTRY: U.S.A.
ZIP: 02110-2804
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30B
CURRENT APPLICATION DATA:
APPLICATION NUMBER: PCT/US95/11511
FILING DATE:

ATTORNEY/AGENT INFORMATION:
NAME: CLARK, PAUL T.
REGISTRATION NUMBER: 30,162
REFERENCE/DOCKET NUMBER: 00786/226001
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617) 542-5070
TELEFAX: (617) 542-8906
TELEX: 200154
INFORMATION FOR SEQ ID NO: 3:
SEQUENCE CHARACTERISTICS:
LENGTH: 34 base pairs
TYPE: nucleic acid
STRANDEDNESS: single
TOPOLOGY: linear
PCT-US95-11511-3

Query Match 0.6%; Score 16; DB 5; Length 34;
Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 225 cttcaacatgtggaag 240
DB 29 CTTCAACATGTGGAAG 14

RESULT 42
US-09-058-969-11
Sequence 11, Application US/09058969A
Patent No. 6228603
GENERAL INFORMATION:
APPLICANT: Reed, John C.
APPLICANT: Deyeraux, Quinn
APPLICANT: Salvesen, Guy S.
APPLICANT: Takahashi, Ryoosuke
APPLICANT: Roy, Natalie
TITLE OF INVENTION: Screening Assays For Agents That Alter Inhibitor of
TITLE OF INVENTION: Apoptosis (IAP) Protein Regulation of Caspase Activity
FILE REFERENCE: LJ 3080
CURRENT APPLICATION NUMBER: US/09/058,969A

;; CURRENT FILING DATE: 1998-04-10
;; EARLIER APPLICATION NUMBER: 08/862,087
;; EARLIER FILING DATE: 1997-05-22
;; NUMBER OF SEQ ID NOS: 12
;; SOFTWARE: Patentln Ver. 2.0
;; SEQ ID NO 11
;; LENGTH: 37
;; TYPE: DNA
;; ORGANISM: Homo sapiens
US-09-058-969-11

Query Match 0.6%; Score 16; DB 4; Length 37;
Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 632 tgaatcctcacaac 647
Db 20 tgaatcctcacaac 35

RESULT 43

US-08-631-200-6
; Sequence 6, Application US/08631200
; Patent No. 5646040
; GENERAL INFORMATION:
; APPLICANT: Kieyn, Patrick W.
; TITLE OF INVENTION: COMPOSITIONS FOR THE TREATMENT AND
; TITLE OF INVENTION: DIAGNOSIS OF BODY WEIGHT DISORDERS, INCLUDING OBESITY
; NUMBER OF SEQUENCES: 59
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Pennie & Edmonds
; STREET: 1155 Avenue of the Americas
; CITY: New York
; STATE: New York
; COUNTRY: U.S.A.
; ZIP: 10036-2711
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patentln Release #1.0, Version #1.30
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/631,200
; FILING DATE: 12-APR-1996
; CLASSIFICATION: 435
; ATTORNEY/AGENT INFORMATION:
; NAME: Coruzzi, Laura A.
; REGISTRATION NUMBER: 30,742
; REFERENCE/DOCKET NUMBER: 7853-057
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (212) 790-9090
; TELEFAX: (212) 869-9741/8864
; TELEX: 66141 PENNIE
; INFORMATION FOR SEQ ID NO: 6:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 39 base pairs
; TYPE: nucleic acid
; STRANDEDNESS: single
; TOPOLOGY: unknown
; MOLECULE TYPE: cDNA
US-08-631-200-6

Query Match 0.6%; Score 16; DB 1; Length 39;
Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 463 gactacatgctatgc 478
Db 15 GACTACATGCTATGC 30

RESULT 44
US-08-829-553-6
; Sequence 6, Application US/08829553
; Patent No. 5817762
; GENERAL INFORMATION:
; APPLICANT: Kieyn, Patrick W.
; TITLE OF INVENTION: COMPOSITIONS FOR THE TREATMENT AND
; TITLE OF INVENTION: DIAGNOSIS OF BODY WEIGHT DISORDERS, INCLUDING OBESITY
; NUMBER OF SEQUENCES: 59
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Pennie & Edmonds
; STREET: 1155 Avenue of the Americas
; CITY: New York
; STATE: New York
; COUNTRY: U.S.A.
; ZIP: 10036-2711
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patentln Release #1.0, Version #1.30
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/829,553
; FILING DATE: 28-MAR-1997
; CLASSIFICATION: 530
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: US 08/631,200
; FILING DATE: 12-APR-1996
; ATTORNEY/AGENT INFORMATION:
; NAME: Coruzzi, Laura A.
; REGISTRATION NUMBER: 30,742
; REFERENCE/DOCKET NUMBER: 7853-057
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (212) 790-9090
; TELEFAX: (212) 869-9741/8864
; TELEX: 66141 PENNIE
; INFORMATION FOR SEQ ID NO: 6:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 39 base pairs
; TYPE: nucleic acid
; STRANDEDNESS: single
; TOPOLOGY: unknown
; MOLECULE TYPE: cDNA
US-08-829-553-6

Query Match 0.6%; Score 16; DB 1; Length 39;
Best Local Similarity 100.0%; Pred. No. 7.4e+02;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 463 gactacatgctatgc 478
Db 15 GACTACATGCTATGC 30

RESULT 45
US-08-922-267A-6
; Sequence 6, Application US/08922267A
; Patent No. 5861239
; GENERAL INFORMATION:
; APPLICANT: Kieyn, Patrick W.
; TITLE OF INVENTION: COMPOSITIONS FOR THE TREATMENT AND
; TITLE OF INVENTION: DIAGNOSIS OF BODY WEIGHT DISORDERS, INCLUDING OBESITY
; NUMBER OF SEQUENCES: 82
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Pennie & Edmonds LLP
; STREET: 1155 Avenue of the Americas
; CITY: New York
; STATE: New York
; COUNTRY: U.S.A.

```

1 ZIP: 10036-2711
2 COMPUTER READABLE FORM:
3 MEDIUM TYPE: Floppy disk
4 COMPUTER: IBM PC compatible
5 OPERATING SYSTEM: PC-DOS/MS-DOS
6 SOFTWARE: Patentin Release #1.0, Version #1.30
7 CURRENT APPLICATION DATA:
8 APPLICATION NUMBER: US/08/922,267A
9 FILING DATE: 2-SEP-1997
10 CLASSIFICATION: 530
11 PRIOR APPLICATION DATA:
12 APPLICATION NUMBER: US 08/929,553
13 FILING DATE: 28-MAR-1997
14 CLASSIFICATION: 530
15 PRIOR APPLICATION DATA:
16 APPLICATION NUMBER: US 08/631,200
17 FILING DATE: 12-APR-1996
18 CLASSIFICATION: 530
19 ATTORNEY/AGENT INFORMATION:
20 NAME: Coruzzi, Laura A.
21 REGISTRATION NUMBER: 30,742
22 REFERENCE/DOCKET NUMBER: 7853-085
23 TELECOMMUNICATION INFORMATION:
24 TELEPHONE: (212) 790-9090
25 TELEFAX: (212) 869-9741/8864
26 TELEX: 66141 PENNIE
27 INFORMATION FOR SEQ ID NO: 6:
28 SEQUENCE CHARACTERISTICS:
29 LENGTH: 39 base pairs
30 TYPE: nucleic acid
31 STRANDEDNESS: single
32 TOPOLOGY: linear
33 MOLECULE TYPE: DNA
34 US-08-922-267A-6
35
36 Query Match 0.68; Score 16; DB 2; Length 39;
37 Best Local Similarity 100.0%; Pred. No. 7.4e+02;
38 Matches 16; Conservative 0; Mismatches 0; Indels 0;
39
40 463 gactcacatcgctatcg 478
41 ||||||||||||||||
42 DB 15 GACTCACATCGCTATCG 30

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Search completed: April 20, 2002, 10:09:55
Job time: 11997 sec

DR WPI; 1999-120935/10

XX Detecting genetic predisposition for body mass disorders - by
PT Identifying allelic variants of a polymorphic region of the SR-BI
PT gene
XX
XX
PS Example 5; Page 72; 102pp; English.
XX
CC A PCR primer pair (see also AAX24561) is designed for the
CC amplification of exon 8 (see AAX24505) of the human SR-BI gene.
CC A C/T polymorphism has been detected at nucleotide 41 of this
CC exon. PCR amplification followed by HaeIII digestion yields
CC 154, 33 and 31 bp products in CC individuals, 154, 64, 33 and 31
CC bp products in CT individuals, and 154 and 64 bp products in TT
CC individuals. The invention is based on the discovery of the
CC genomic structure of the human SR-BI gene (see AAX24498-509) and on
CC the identification of polymorphic regions within the gene which are
CC associated with abnormal body mass index (BMI) and abnormal
CC lipoprotein levels and hence with disorders such as obesity,
CC cachexia, cardiovascular disorders and gallstone formation. The
CC invention provides methods for determining whether a subject has,
CC or is at risk of developing, a disease associated with a specific
CC allele of a polymorphic region of an SR-BI gene. Kits comprising
CC the relevant probe or primer are claimed.
XX
SQ Sequence 34 BP; 4 A; 15 C; 3 G; 12 T; 0 other;

Query Match 1.2%; Score 31; DB 20; Length 34;
Best Local Similarity 100.0%; Pred. No. 0.0013;
Matches 31; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1085 cctgttctctccatcctcattcctca 1115
Db 1 cctgttctctccatcctcattcctca 31
|||||

RESULT 2
AAX24652
ID AAX24652 standard; DNA; 34 BP.
XX
AC AAX24652;
XX
DT 21-JUN-1999 (first entry)
XX
DE Human SR-BI gene exon 8 PCR primer.
XX
KW SR-BI; human; polymorphism; cardiovascular disorder; ischaemia;
KW restenosis; congestive heart failure; atherosclerosis; cholesterol;
KW low density lipoprotein; LDL; high density lipoprotein; HDL;
KW diagnosis; body mass index; obesity; cachexia; gallstone; PCR;
KW primer; ss.
XX
OS Synthetic.
OS Homo sapiens.
XX
PN WO9902736-A2.
XX
PD 21-JAN-1999.
XX
PE 10-JUL-1998; 98WO-US14359.
XX
PR 27-FEB-1998; 98US-0032894.
XX
PR 10-JUL-1997; 97US-0890980.
XX
PA (MILL-) MILLENNIUM PHARM INC.
XX
PI Action SL;
XX
DR WPI; 1999-120936/10.
XX
PT New nucleic acids comprising intronic sequence of a human scavenger
PT receptor-BI (SR-BI) gene - useful for prognosis, diagnosis and
PT treatment of SR-BI associated diseases or conditions

XX Claim 10; Page 71; 103pp; English.
PS
XX
CC A PCR primer pair (see also AAX24597) is designed for the
CC amplification of exon 8 (see AAX24597) of the human SR-BI gene.
CC A C/T polymorphism has been detected at nucleotide 41 of this
CC exon. PCR amplification followed by HaeIII digestion yields
CC 154, 33 and 31 bp products in CC individuals, 154, 64, 33 and 31
CC bp products in CT individuals, and 154 and 64 bp products in TT
CC individuals. The invention is based on the discovery of the
CC genomic structure of the human SR-BI gene (see AAX24590-601) and on
CC the identification of polymorphic regions within the gene which are
CC associated with abnormal body mass index (BMI) and abnormal
CC lipoprotein levels and hence with disorders such as obesity,
CC cachexia, cardiovascular disorders and gallstone formation. The
CC invention provides methods for determining whether a subject has,
CC or is at risk of developing, a disease associated with a specific
CC allele of a polymorphic region of an SR-BI gene. Kits comprising
CC the relevant probe or primer are claimed.
XX
SQ Sequence 34 BP; 4 A; 15 C; 3 G; 12 T; 0 other;

Query Match 1.2%; Score 31; DB 20; Length 34;
Best Local Similarity 100.0%; Pred. No. 0.0013;
Matches 31; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1085 cctgttctctccatcctcattcctca 1115
Db 1 cctgttctctccatcctcattcctca 31
|||||

RESULT 3
AA21075/C
ID AA21075 standard; DNA; 30 BP.
XX
AC AA21075;
XX
DT 18-NOV-1999 (first entry)
XX
DE Human cell-surface HDL receptor CLA-1 probe.
XX
KW LDL receptor; low density lipoprotein; steroid receptor element;
KW caveolin; SRE; regulation; cell cycle; cholesterol; mitosis;
KW cell division; anti-mitotic; inhibition; growth; proliferation;
KW cancer; restenosis; atherosclerosis; heart disease; detection;
KW lipid processing; diabetes; thyroid hormone deficiency; renal failure;
KW inherited hyperlipidaemia; probe; ss.
XX
OS Synthetic.
OS Homo sapiens.
XX
PN WO9946592-A1.
XX
PD 16-SEP-1999.
XX
PE 08-MAR-1999; 99WO-US05146.
XX
PR 09-MAR-1998; 98US-0077351.
XX
PA (REGC) UNIV CALIFORNIA.
XX
PI Fielding CJ, Fielding PE;
XX
DR WPI; 1999-551504/46.
XX
PT Detection of anti-mitotic agents for use in inhibiting the growth or
PT proliferation of cells, e.g. in cancers or restenosis.
XX
PS Example 5; Page 92; 135pp; English.
XX
CC A method has been developed for identifying anti-mitotic agents by
CC detecting effects on cholesterol influx or efflux in cells or using a

Query Match

Best Local Similarity 0.9%; Score 24; DB 20; Length 24;
Pred. No. 1.3;
Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 67 gacatggcgtctccgcaagcg 90
|||||
DB 24 GACATGGCGTCTCCGCCAAGCG 1

RESULT 6

AA24603/c
ID AAX24603 standard; DNA; 24 BP.

AC AAX24603;

DT 21-JUN-1999 (first entry)

DE Human SR-BI gene exon 1 primer 3e16srbl.

XX SR-BI; human; polymorphism; cardiovascular disorder; ischaemia;
XX restenosis; congestive heart failure; atherosclerosis; cholesterol;
XX low density lipoprotein; LDL; high density lipoprotein; HDL;
XX diagnosis; body mass index; obesity; cachexia; gallstone; PCR;
XX primer; ss.

XX Synthetic.

OS Homo sapiens.

PN WO9902736-A2.

PD 21-JAN-1999.

PF 10-JUL-1998; 98WO-US14359.

PR 27-FEB-1998; 98US-0032894.

PR 10-JUL-1997; 97US-0890980.

PA (MILL-) MILLENNIUM PHARM INC.

PI Action SL;

DR WPI; 1999-120936/10.

XX New nucleic acids comprising intronic sequence of a human scavenger
PT receptor-BI (SR-BI) gene - useful for prognosis, diagnosis and
PT treatment of SR-BI associated diseases or conditions

PS Claim 10; Page 66; 103pp; English.

XX Primer 3e16srbl is used with primer 5e16srbl (see AAX24602) in the
CC PCR amplification of exon 1 (see AAX24590) of the human SR-BI gene.
CC The invention is based on the discovery of the genomic structure of
CC the human SR-BI gene (see AAX24590-601) and on the identification of
CC polymorphic regions within the gene which are associated with
CC abnormal body mass index (BMI) and abnormal lipoprotein levels and
CC hence with disorders such as obesity, cachexia, cardiovascular
CC disorders and gallstone formation. Claimed primers (see AAX24602-25)
CC are used for the amplification of the exons, introns and promoter
CC region of the SR-BI gene for detection of polymorphisms and
CC mutations. The invention provides methods for determining whether
CC a subject has, or is at risk of developing, a disease associated
CC with a specific allele of a polymorphic region of an SR-BI gene.
CC Kits comprising the relevant probe or primer are claimed.

SO Sequence 24 BP; 3 A; 8 C; 8 G; 5 T; 0 other;

Query Match 0.9%; Score 24; DB 20; Length 24;

Best Local Similarity 100.0%; Pred. No. 1.3;
Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 67 gacatggcgtctccgcaagcg 90

DB 24 GACATGGCGTCTCCGCCAAGCG 1
|||||

RESULT 7
AAX24557/c
ID AAX24557 standard; DNA; 23 BP.

AC AAX24557;

DT 21-JUN-1999 (first entry)

DE Human SR-BI gene exon 1 PCR primer.

XX SR-BI; human; polymorphism; cardiovascular disorder; ischaemia;
XX restenosis; congestive heart failure; atherosclerosis; cholesterol;
XX low density lipoprotein; LDL; high density lipoprotein; HDL;
XX diagnosis; body mass index; obesity; cachexia; gallstone; PCR;
XX primer; ss.

XX Synthetic.

OS Homo sapiens.

PN WO9902735-A2.

PD 21-JAN-1999.

PF 10-JUL-1998; 98WO-US14354.

PR 27-FEB-1998; 98US-0031626.

PR 10-JUL-1997; 97US-0890979.

PA (MILL-) MILLENNIUM PHARM INC.
(UYTU-) UNIV TUPITS.

PI Action SL; Ordovas JM;

DR WPI; 1999-120935/10;

XX Detecting genetic predisposition for body mass disorders - by
PT identifying allelic variants of a polymorphic region of the SR-BI
PT gene

PS Example 5; Page 72; 102pp; English.

XX A PCR primer pair (see also: AAX24556) is designed for the
CC amplification of exon 1 (see AAX24498) of the human SR-BI gene.
CC A G/A polymorphism has been detected at nucleotide 146 of this
CC exon. PCR amplification followed by AluI digestion yields
CC a 263 bp product in GG individuals, 263, 192 and 71 bp products
CC in GA individuals, and 192 and 71 bp products in AA individuals.
CC The invention is based on the discovery of the genomic structure of
CC the human SR-BI gene (see AAX24498-509) and on the identification of
CC polymorphic regions within the gene which are associated with
CC abnormal body mass index (BMI) and abnormal lipoprotein levels and
CC hence with disorders such as obesity, cachexia, cardiovascular
CC disorders and gallstone formation. The invention provides methods
CC for determining whether a subject has, or is at risk of developing,
CC a disease associated with a specific allele of a polymorphic region
CC of an SR-BI gene. Kits comprising the relevant probe or primer are
CC claimed.

SO Sequence 23 BP; 6 A; 10 C; 6 G; 1 T; 0 other;

Query Match 0.9%; Score 23; DB 20; Length 23;

Best Local Similarity 100.0%; Pred. No. 3.6;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 123 gctactgtgcgtctgtgagcg 145
|||||

DB 23 GCTACTGTGCGTCTGTGAGCG 1

```

RESULT 8
AC AAX24543/C
ID AAX24649 standard; DNA: 23 BP.
XX
AC AAX24649;
XX
DT 21-JUN-1999 (first entry)
XX
DE Human SR-BI gene exon 1 PCR primer.
XX
KM SR-BI; human; polymorphism; cardiovascular disorder; ischaemia;
KM restenosis; congestive heart failure; atherosclerosis; cholesterol;
KM low density lipoprotein; LDL; high density lipoprotein; HDL;
KM diagnosis; body mass index; obesity; cachexia; gallstone; PCR;
KM primer; ss.
XX
OS Synthetic.
OS Homo sapiens.
XX
PN MO9902736-A2.
XX
PD 21-JAN-1999.
XX
PF 10-JUL-1998; 98WO-US14359.
XX
PR 27-FEB-1998; 98US-0032894.
PR 10-JUL-1997; 97US-0890980.
XX
PA (MILL-) MILLENNIUM PHARM INC.
XX
PI Action SL;
XX
DR WPI; 1999-120936/10.
XX
PT New nucleic acids comprising intronic sequence of a human scavenger
PT receptor-BI (SR-BI) gene - useful for prognosis, diagnosis and
PT treatment of SR-BI associated diseases or conditions
XX
PS Claim 10; Page 71; 103pp; English.
XX
XX A PCR primer pair (see also AAX24648) is designed for the
CC amplification of exon 1 (see AAX24590) of the human SR-BI gene.
CC A G/A polymorphism has been detected at nucleotide 146 of this
CC exon. PCR amplification followed by AluI digestion yields
CC a 263 bp product in Gg individuals, 263, 192 and 71 bp products
CC in GA individuals, and 192 and 71 bp products in AA individuals.
CC The invention is based on the discovery of the genomic structure of
CC the human SR-BI gene (see AAX24590-601) and on the identification of
CC polymorphic regions within the gene which are associated with
CC abnormal body mass index (BMI) and abnormal lipoprotein levels and
CC hence with disorders such as obesity, cachexia, cardiovascular
CC disorders and gallstone formation. The invention provides methods
CC for determining whether a subject has, or is at risk of developing,
CC a disease associated with a specific allele of a polymorphic region
CC of an SR-BI gene. Kits comprising the relevant probe or primer are
CC claimed.
XX
SQ Sequence 23 BP; 6 A; 10 C; 6 G; 1 T; 0 other;
XX
Query Match 0.9%; Score 23; DB 20; Length 23;
Best Local Similarity 100.0%; Pred. No. 3; 6; Mismatches 0;
Matches 23; Conservative 0; Indels 0; Gaps 0;
OY 123 gctactgtgcgtgctgtgctgagcg 145
DB 23 GCTACTGTGCGCTGTGCTGCGCG 1
XX
RESULT 9
AC AAX24543/C
ID AAX24543 standard; DNA: 31 BP.
XX

```

```

XX AAX24543;
AC
XX
DT 21-JUN-1999 (first entry)
XX
DE Human SR-BI gene exon 8 probe.
XX
KM SR-BI; human; polymorphism; cardiovascular disorder; ischaemia;
KM restenosis; congestive heart failure; atherosclerosis; cholesterol;
KM low density lipoprotein; LDL; high density lipoprotein; HDL;
KM diagnosis; body mass index; obesity; cachexia; gallstone;
KM probe; hybridisation; ss.
XX
OS Synthetic.
OS Homo sapiens.
XX
PN MO9902735-A2.
XX
PD 21-JAN-1999.
XX
PF 10-JUL-1998; 98WO-US14354.
XX
PR 27-FEB-1998; 98US-0031626.
PR 10-JUL-1997; 97US-0890979.
XX
PA (MILL-) MILLENNIUM PHARM INC.
PA (UTU-) UNIV TUFTS.
XX
PI Action SL; Ordovas JM;
XX
DR WPI; 1999-120935/10.
XX
PT Detecting genetic predisposition for body mass disorders - by
PT identifying allelic variants of a polymorphic region of the SR-BI
PT gene
XX
PS Example 2; Page 33; 102pp; English.
XX
XX This probe is designed to detect a C/T polymorphism located at
CC nucleotide 41 of exon 8 of the human SR-BI gene (see AAX24536).
CC It hybridises specifically to a nucleotide sequence wherein
CC nucleotide 41 is cytidine. The invention is based on the
CC discovery of the genomic structure of the human SR-BI gene (see
CC AAX24498-509) and on the identification of polymorphic regions within
CC the gene which are associated with abnormal body mass index (BMI)
CC and abnormal lipoprotein levels and hence with disorders such as
CC obesity, cachexia, cardiovascular disorders and gallstone formation.
CC The invention provides methods for determining whether a subject
CC has, or is at risk of developing, a disease associated with a
CC specific allele of a polymorphic region of an SR-BI gene. Kits
CC comprising the relevant probe or primer are claimed.
XX
SQ Sequence 31 BP; 7 A; 6 C; 12 G; 6 T; 0 other;
XX
Query Match 0.9%; Score 23; DB 20; Length 31;
Best Local Similarity 100.0%; Pred. No. 3; 5; Mismatches 0;
Matches 23; Conservative 0; Indels 0; Gaps 0;
OY 1112 tcaacgcccagccgcttgcga 1134
DB 23 TCACGCCGACCCGCTGTGCGCA 1
XX
RESULT 10
AC AAX24545
ID AAX24545 standard; DNA: 31 BP.
XX
AC AAX24545;
XX
DT 21-JUN-1999 (first entry)
XX
DE Human SR-BI gene exon 8 probe.
XX

```

XX SR-BI; human; polymorphism; cardiovascular disorder; ischemia;
 KW restenosis; congestive heart failure; atherosclerosis; cholesterol;
 KW low density lipoprotein; LDL; high density lipoprotein; HDL;
 KW diagnosis; body mass index; obesity; cachexia; gallstone;
 KW probe; hybridisation; ss.
 OS Synthetic.
 OS Homo sapiens.
 XX WO9902735-A2.
 PN 21-JAN-1999.
 XX 10-JUL-1998; 98WO-US14354.
 PF 27-FEB-1998; 98US-0031626.
 PR 10-JUL-1997; 97US-0890979.
 XX (MILL-) MILLENNIUM PHARM INC.
 PA (UYTU-) UNIV TUFTS.
 PI Acton ST, Ordovas JM;
 XX WPI; 1999-120935/10.
 DR
 XX
 PT Detecting genetic predisposition for body mass disorders - by
 PT identifying allele variants of a polymorphic region of the SR-BI
 gene
 PS Example 2; Page 33; 102pp; English.
 XX This probe is designed to detect a C/T polymorphism located at
 CC nucleotide 41 of exon 8 of the human SR-BI gene (see AAX24636).
 CC It hybridises specifically to the complement of a nucleotide
 CC sequence wherein nucleotide 41 is cytidine. The invention is
 CC based on the discovery of the genomic structure of the human SR-BI
 CC gene (see AAX24498-509) and on the identification of polymorphic
 CC regions within the gene which are associated with abnormal body
 CC mass index (BMI) and abnormal lipoprotein levels and hence with
 CC disorders such as obesity, cachexia, cardiovascular disorders and
 CC gallstone formation. The invention provides methods for
 CC determining whether a subject has, or is at risk of developing, a
 CC disease associated with a specific allele of a polymorphic region
 CC of an SR-BI gene. Kits comprising the relevant probe or primer are
 CC claimed.
 XX Sequence 31 BP; 6 A; 12 C; 6 G; 7 T; 0 other;
 SQ

Query Match 0.9%; Score 23; DB 20; Length 31;
 Best Local Similarity 100.0%; Pred. No. 3.5;
 Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1112 tcaacgccgacccggtctgca 1134
 ||||||||||||||||||||
 Db 9 tcaacgccgacccggtctgca 31

RESULT 11
 AAX24635/C
 ID AAX24635 standard; DNA; 31 BP.
 XX
 AC AAX24635;
 XX
 DT 21-JUN-1999 (first entry)
 XX
 DE Human SR-BI gene exon 8 probe.
 XX
 KW SR-BI; human; polymorphism; cardiovascular disorder; ischemia;
 KW restenosis; congestive heart failure; atherosclerosis; cholesterol;
 KW low density lipoprotein; LDL; high density lipoprotein; HDL;
 KW diagnosis; body mass index; obesity; cachexia; gallstone;
 KW probe; hybridisation; ss.
 OS Synthetic.
 OS Homo sapiens.
 XX WO9902736-A2.
 PN 21-JAN-1999.
 XX 10-JUL-1998; 98WO-US14359.
 PF 27-FEB-1998; 98US-0032894.
 PR 10-JUL-1997; 97US-0890980.
 XX (MILL-) MILLENNIUM PHARM INC.
 PA Acton ST;
 XX WPI; 1999-120936/10.
 DR
 XX
 PT New nucleic acids comprising intronic sequence of a human scavenger
 PT receptor-BI (SR-BI) gene - useful for prognosis, diagnosis and
 PT treatment of SR-BI associated diseases or conditions
 PS Claim 36; Page 32; 103pp; English.
 XX This probe is designed to detect a C/T polymorphism located at
 CC nucleotide 41 of exon 8 of the human SR-BI gene (see AAX24628).
 CC It hybridises specifically to a nucleotide sequence wherein
 CC nucleotide 41 of exon 8 is cytidine. The invention is based on
 CC the discovery of the genomic structure of the human SR-BI gene (see
 CC AAX24590-601) and on the identification of polymorphic regions within
 CC the gene which are associated with abnormal body mass index (BMI)
 CC and abnormal lipoprotein levels and hence with disorders such as
 CC obesity, cachexia, cardiovascular disorders and gallstone formation.
 CC The invention provides methods for determining whether a subject
 CC has, or is at risk of developing, a disease associated with a
 CC specific allele of a polymorphic region of an SR-BI gene. Kits
 CC comprising the relevant probe or primer are claimed.
 XX Sequence 31 BP; 7 A; 6 C; 12 G; 6 T; 0 other;
 SQ

KW probe; hybridisation; ss.
 XX
 OS Synthetic.
 OS Homo sapiens.
 XX WO9902736-A2.
 PN 21-JAN-1999.
 XX 10-JUL-1998; 98WO-US14359.
 PF 27-FEB-1998; 98US-0032894.
 PR 10-JUL-1997; 97US-0890980.
 XX (MILL-) MILLENNIUM PHARM INC.
 PA Acton ST;
 XX WPI; 1999-120936/10.
 DR
 XX
 PT New nucleic acids comprising intronic sequence of a human scavenger
 PT receptor-BI (SR-BI) gene - useful for prognosis, diagnosis and
 PT treatment of SR-BI associated diseases or conditions
 PS Claim 36; Page 32; 103pp; English.
 XX This probe is designed to detect a C/T polymorphism located at
 CC nucleotide 41 of exon 8 of the human SR-BI gene (see AAX24628).
 CC It hybridises specifically to a nucleotide sequence wherein
 CC nucleotide 41 of exon 8 is cytidine. The invention is based on
 CC the discovery of the genomic structure of the human SR-BI gene (see
 CC AAX24590-601) and on the identification of polymorphic regions within
 CC the gene which are associated with abnormal body mass index (BMI)
 CC and abnormal lipoprotein levels and hence with disorders such as
 CC obesity, cachexia, cardiovascular disorders and gallstone formation.
 CC The invention provides methods for determining whether a subject
 CC has, or is at risk of developing, a disease associated with a
 CC specific allele of a polymorphic region of an SR-BI gene. Kits
 CC comprising the relevant probe or primer are claimed.
 XX Sequence 31 BP; 7 A; 6 C; 12 G; 6 T; 0 other;
 SQ

Query Match 0.9%; Score 23; DB 20; Length 31;
 Best Local Similarity 100.0%; Pred. No. 3.5;
 Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1112 tcaacgccgacccggtctgca 1134
 ||||||||||||||||||||
 Db 23 TCAACGCCGACCCGCTCTGCA 1

RESULT 12
 AAX24637
 ID AAX24637 standard; DNA; 31 BP.
 XX
 AC AAX24637;
 XX
 DT 21-JUN-1999 (first entry)
 XX
 DE Human SR-BI gene exon 8 probe.
 XX
 KW SR-BI; human; polymorphism; cardiovascular disorder; ischemia;
 KW restenosis; congestive heart failure; atherosclerosis; cholesterol;
 KW low density lipoprotein; LDL; high density lipoprotein; HDL;
 KW diagnosis; body mass index; obesity; cachexia; gallstone;
 KW probe; hybridisation; ss.
 OS Synthetic.
 OS Homo sapiens.
 XX WO9902736-A2.
 PN 21-JAN-1999.
 XX 10-JUL-1998; 98WO-US14359.
 PF 27-FEB-1998; 98US-0032894.
 PR 10-JUL-1997; 97US-0890980.
 XX (MILL-) MILLENNIUM PHARM INC.
 PA Acton ST;
 XX WPI; 1999-120936/10.
 DR
 XX
 PT New nucleic acids comprising intronic sequence of a human scavenger
 PT receptor-BI (SR-BI) gene - useful for prognosis, diagnosis and
 PT treatment of SR-BI associated diseases or conditions
 PS Claim 36; Page 32; 103pp; English.
 XX This probe is designed to detect a C/T polymorphism located at
 CC nucleotide 41 of exon 8 of the human SR-BI gene (see AAX24628).
 CC It hybridises specifically to a nucleotide sequence wherein
 CC nucleotide 41 of exon 8 is cytidine. The invention is based on
 CC the discovery of the genomic structure of the human SR-BI gene (see
 CC AAX24590-601) and on the identification of polymorphic regions within
 CC the gene which are associated with abnormal body mass index (BMI)
 CC and abnormal lipoprotein levels and hence with disorders such as
 CC obesity, cachexia, cardiovascular disorders and gallstone formation.
 CC The invention provides methods for determining whether a subject
 CC has, or is at risk of developing, a disease associated with a
 CC specific allele of a polymorphic region of an SR-BI gene. Kits
 CC comprising the relevant probe or primer are claimed.
 XX Sequence 31 BP; 7 A; 6 C; 12 G; 6 T; 0 other;
 SQ

PD 21-JAN-1999.
 XX 10-JUL-1998; 98WO-US14359.
 XX 27-FEB-1998; 98US-0032894.
 PR 10-JUL-1997; 97US-0890980.
 XX (MILL-) MILLENNIUM PHARM INC.
 XX Acton SL;
 PI WPI; 1999-120936/10.
 DR New nucleic acids comprising intronic sequence of a human scavenger
 XX receptor-BI (SR-BI) gene - useful for prognosis, diagnosis and
 PT treatment of SR-BI associated diseases or conditions
 XX
 PS Claim 36; Page 32; 103pp; English.
 CC This probe is designed to detect a C/T polymorphism located at
 CC nucleotide 41 of exon 8 of the human SR-BI gene (see AAX24628).
 CC It hybridises specifically to the complement of a sequence wherein
 CC nucleotide 41 of exon 8 is cytidine. The invention is based on
 CC the discovery of the genomic structure of the human SR-BI gene (see
 CC AAX24590-601) and on the identification of polymorphic regions within
 CC the gene which are associated with abnormal body mass index (BMI)
 CC and abnormal lipoprotein levels and hence with disorders such as
 CC obesity, cachexia, cardiovascular disorders and gallstone formation.
 CC The invention provides methods for determining whether a subject
 CC has, or is at risk of developing, a disease associated with a
 CC specific allele of a polymorphic region of an SR-BI gene. Kits
 CC comprising the relevant probe or primer are claimed.
 XX
 SO Sequence 31 BP; 6 A; 12 C; 6 G; 7 T; 0 other;

Query Match 0.9%; Score 23; DB 20; Length 31;
 Best Local Similarity 100.0%; Pred. No. 3.5;
 Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaagcgagccgggtctgcga 1134
 |||||||
 DB 9 tcaagcgagccgggtctgcga 31

RESULT 13
 AAQ75789/C
 ID AAQ75789 standard; DNA; 21 BP.
 XX
 AC AAQ75789;
 XX
 DT 04-AUG-1995 (first entry)
 XX
 DE Reverse transcription primer used in cDNA analysis technique.
 XX
 XX Analysis; gene expression; reverse transcription; primer; cDNA;
 KW aggregate; restriction enzyme; ss.
 XX
 OS Synthetic.
 XX
 PN JP06303997-A.
 XX
 PD 01-NOV-1994.
 XX
 PF 16-APR-1993; 93JP-0112515.
 XX
 PR 16-APR-1993; 93JP-0112515.
 XX
 PA (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.
 XX
 DR WPI; 1995-018287/03.
 XX
 PT Analysis of cDNA and gene expression - by amplification of mRNA

PT followed by digestion with restriction enzymes
 XX
 PS Disclosure; Page 9; 11pp; Japanese.
 XX
 CC A method for the analysis of cDNA comprises (a) preparing an
 CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs
 CC and a plural type of labelled reverse transcription primers
 CC (GENSEQ files AAQ75547-075798) and using the aggregate of mRNAs as the
 CC template for each reverse transcription primer; (b) digesting each of
 CC the prepared aggregates of the double-stranded cDNAs with restriction
 CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
 CC separate lanes. The method can be used to analyse gene expression
 CC rapidly and easily.
 XX
 SO Sequence 21 BP; 1 A; 2 C; 0 G; 18 T; 0 other;

Query Match 0.7%; Score 18; DB 16; Length 21;
 Best Local Similarity 100.0%; Pred. No. 5.1e+02;
 Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2549 atgagaaaaaataaaaaa 2566
 |||||||
 DB 21 ATCGAAAAAATAAAAAA 4

RESULT 14
 AAV11549
 ID AAV11549 standard; cDNA; 36 BP.
 XX
 AC AAV11549;
 XX
 DT 14-SEP-1998 (first entry)
 XX
 DE Human SR-BI gene PCR primer SRB1 5'1387.
 XX
 KW Lipid metabolic pathway; h-LMP-1 gene; cardiovascular disease;
 KW atherosclerosis; biliary tract disorder; gall stone; therapy;
 KW diagnosis; human; SR-BI; PCR; primer; ss.
 XX
 OS Synthetic.
 OS Homo sapiens.
 OS
 PN WO9809979-A1.
 XX
 PD 12-MAR-1998.
 XX
 PF 28-AUG-1997; 97WO-US15195.
 XX
 PR 04-SEP-1996; 96US-0707399.
 XX
 PA (MILL-) MILLENNIUM PHARM INC.
 XX
 PI Acton S; Gimeno CJ;
 XX
 DR WPI; 1998-193545/17.
 XX
 PT DNA encoding lipid metabolic pathway polypeptide(s) - useful for
 PT treatment of cardiovascular disease or modulation of lipid uptake or
 PT metabolism
 XX
 PS Example 1; Page 84; 102pp; English.
 XX
 CC PCR primer SRB1 5'1387 was used with primer SRB1 3'1528r to amplify
 CC cDNA encoding amino acids 463-509 (i.e. the cytoplasmic domain) of
 CC human SR-BI. Restriction endonuclease EcoRI and BamHI sites were
 CC engineered into the oligonucleotides to allow the cloning of the
 CC SR-BI cytoplasmic domain into two-hybrid system DNA-binding domain
 CC fusion vector pGBT9. This was used to identify a novel gene (see
 CC AAV11547) coding for human lipid metabolic pathway (LMP) protein (see
 CC AAV58888). LMP nucleic acids and polypeptides are useful for
 CC developing methods for treatment of cardiovascular diseases or
 CC for modulating lipid uptake or metabolism.

XX Sequence 36 BP; 10 A; 9 C; 10 G; 7 T; 0 other;

Query Match 0.7%; Score 18; DB 19; Length 36;
Best Local Similarity 100.0%; Pred. No. 4.9e+02;
Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1456 caaatccggagcccaagag.1473
DB 19 caaatccggagcccaagag 36

RESULT 15
ID AAH20331/C
AAH20331; DNA: 40 BP.

AC AAH20331;
XX
DT 01-AUG-2001 (first entry)

DE Rabies virus glycoprotein specific primer RGP/R797 SEQ ID 12.

KW Primer; solid phase amplification of DNA template; SPADT; detection; RGP;
KM genomic scanning; bacterial diagnostic; rabies virus glycoprotein; ss.

OS Rabies virus.
OS Synthetic.

PN US6221635-B1.

PD 24-APR-2001.

PF 06-MAY-1999; 99US-0306290.

PR 06-MAY-1999; 99US-0306290.

PA (WIS-) WISTAR INST.

PI Rovera G, Mukhopadhyay S;

DR WPI; 2001-315577/33.

PT Detecting the presence of a specific nucleic acid in a sample
PT containing DNA, useful in scanning large genomic fragments for the
PT presence of genes or gene families, comprises performing solid phase
PT amplification of DNA template

XX Example 1; Column 22; 49pp; English.

CC This invention relates to a method for detecting the presence of a
CC specific nucleic acid in a sample containing DNA. The method comprises
CC performing solid phase amplification of DNA template (SPADT). 5' and 3'
CC primers are irreversibly bound to a solid support, and the DNA from a
CC sample is absorbed and reversibly bound, incubated under amplification
CC reaction conditions and the presence of the specific target DNA is
CC detected. The method is useful for detecting the presence of a specific
CC nucleic acid (e.g. bacterial, viral or parasitic DNA) in a sample or in a
CC cell. SPADT may be used for scanning large genomic fragments for the
CC presence of genes or gene families; or for bacterial diagnostics by
CC examining the ribosomal RNA genes; or for viral diagnostics by scanning
CC for the presence of viral nucleic acid sequences in a sample. SPADT may
CC also be used in forensic medicine by detecting and identifying species
CC specific sequences or for the presence of major histocompatibility
CC complex. The present sequence represents a primer specific for the rabies
CC virus glycoprotein gene (RGP). The primer is used in an example
CC illustrating the method of the invention.

SO Sequence 40 BP; 6 A; 7 C; 2 G; 25 T; 0 other;

Query Match 0.7%; Score 18; DB 22; Length 40;
Best Local Similarity 100.0%; Pred. No. 4.9e+02;

Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2549 atggaataaaaaaa 2566
DB 24 ATGGAATAAAAAAAA 7

RESULT 16
ID AAX69803/C
AAX69803; RNA: 17 BP.

AC AAX69803;
XX
DT 28-JUL-1999 (first entry)

DE Human flt1 VEGF receptor hammethead ribozyme substrate #1098.

KW Vascular endothelial growth factor receptor; VEGF receptor; flt-1;
KM flk-1; KDR; hammethead ribozyme; hairpin ribozyme; cleavage;
KM tumour angiogenesis; psoriasis; rheumatoid arthritis; ocular disease;
KM flk-like tyrosine kinase 1; kinase insert domain containing receptor;
KM foetal liver kinase 1; ss.

XX Homo sapiens.
OS
XX WO9715662-A2.

PD 01-MAY-1997;
XX 25-OCT-1996; 96WO-US17480.

PF 11-JAN-1996; 96US-0584040.

PR 26-OCT-1995; 95US-0005974.

PA (CHIR) CHIRON CORP.
PA (RIBO-) RIBOZYME PHARM INC.

PI Escobedo J, McSwiggen J, Pavco P, Stinchcomb D;

DR WPI; 1997-259017/23.

PT Nucleic acid molecule modulating VEGF receptor(s) gene expression or
PT mRNA stability - useful for treating e.g. tumour angiogenesis,
PT psoriasis, rheumatoid arthritis, etc., in a human patient

PS Claim 4; Page 79; 218pp; English.

XX The present invention describes nucleic acid molecules which modulate
CC the synthesis, expression and/or stability of a mRNA encoding 1 or more
CC receptors of vascular endothelial growth factor (VEGF). A patient
CC (preferably human) having a condition associated with the level of the
CC flk-like tyrosine kinase 1 (flt-1), kinase insert domain containing
CC receptor (KDR) and/or foetal liver kinase 1 (flk-1) (e.g. tumour
CC angiogenesis, ocular diseases, psoriasis and rheumatoid arthritis) can
CC be treated by administering the nucleic acid molecule or the expression
CC vector to the patient. AAX67275 to AAX75752 represent specific examples
CC of nucleic acid molecules from the present invention.

SO Sequence 17 BP; 1 A; 2 C; 0 G; 14 U; 0 other;

Query Match 0.7%; Score 17; DB 18; Length 17;
Best Local Similarity 100.0%; Pred. No. 1.4e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2550 tggaaaaaa 2566
DB 17 TGGAAAAAAA 1

RESULT 17
ID AAG75604/C
AAG75604 standard; DNA: 20 BP.


```
XX AA075604;
AC
XX
DT 04-AUG-1995 (first entry)
DE Reverse transcription primer used in cDNA analysis technique.
XX
XX Analysis: gene expression; reverse transcription; primer; cDNA;
XX aggregate; restriction enzyme; ss.
XX
OS Synthetic.
XX
XX JP06303997-A.
XX
XX 01-NOV-1994.
XX
XX 16-APR-1993; 93JP-0112515.
XX
XX 16-APR-1993; 93JP-0112515.
XX
XX (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.
XX
XX WPI: 1995-018287/03.
XX
XX Analysis of cDNA and gene expression - by amplification of mRNA
XX followed by digestion with restriction enzymes
XX
XX Disclosure; Page 5; 11pp; Japanese.
XX
XX A method for the analysis of cDNA comprises (a) preparing an
XX aggregate of double-stranded cDNAs by using an aggregate of mRNAs
XX and a plural type of labelled reverse transcription primers
XX (GENSEQ files AA075547-075798) and using the aggregate of mRNAs as the
XX template for each reverse transcription primer; (b) digesting each of
XX the prepared aggregates of the double-stranded cDNAs with restriction
XX enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
XX separate lanes. The method can be used to analyse gene expression
XX rapidly and easily.
XX
XX Sequence 20 BP; 1 A; 2 C; 0 G; 17 T; 0 other:
XX
Query Match 0.7%; Score 17; DB 16; Length 20;
Best Local Similarity 100.0%; Pred. No. 1.4e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 2550 tggagaaaaa 2566
Db 20 TCGAAAAA 4
RESULT 18
AA075604/C
ID AAX24542 standard; DNA; 20 BP.
XX
XX AAX24542;
AC
XX
XX 21-JUN-1999 (first entry)
DT
XX
XX Human SR-BI gene exon 8 probe.
DE
XX
XX SR-BI: human; polymorphism; cardiovascular disorder; ischaemia;
XX restenosis; congestive heart failure; atherosclerosis; cholesterol;
XX low density lipoprotein; LDL; high density lipoprotein; HDL;
XX diagnosis; body mass index; obesity; cachexia; gallstone;
XX probe; hybridisation; ss.
XX
XX Synthetic.
XX Homo sapiens.
XX
XX WO9902735-A2.
XX
XX 21-JAN-1999.
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XX
XX 10-JUL-1998; 98MO-US14354.
XX
XX 27-FEB-1998; 98US-0031626.
XX
XX 10-JUL-1997; 97US-0890979.
XX
XX (MILL-) MILLENNIUM PHARM INC.
XX (UYTU-) UNIV TUFTS.
XX
XX Acton ST, Ordovas JM.
XX
XX WPI: 1999-120935/10.
XX
XX Detecting genetic predisposition for body mass disorders - by
XX identifying allelic variants of a polymorphic region of the SR-BI
XX gene
XX
XX Example 2; Page 33; 102pp; English.
XX
XX This probe is designed to detect a C/T polymorphism located at
XX nucleotide 41 of exon 8 of the human SR-BI gene (see AAX24536).
XX It hybridises specifically to a nucleotide sequence wherein
XX nucleotide 41 is cytidine. The invention is based on the
XX discovery of the genomic structure of the human SR-BI gene (see
XX AAX24498-509) and on the identification of polymorphic regions within
XX the gene which are associated with abnormal body mass index (BMI)
XX and abnormal lipoprotein levels and hence with disorders such as
XX obesity, cachexia, cardiovascular disorders and gallstone formation.
XX The invention provides methods for determining whether a subject
XX has, or is at risk of developing, a disease associated with a
XX specific allele of a polymorphic region of an SR-BI gene. Kits
XX comprising the relevant probe or primer are claimed.
XX
XX Sequence 20 BP; 4 A; 4 C; 8 G; 4 T; 0 other:
XX
Query Match 0.7%; Score 17; DB 20; Length 20;
Best Local Similarity 100.0%; Pred. No. 1.4e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1112 tcaagcgagcccggtt 1128
Db 17 TCAAGCGGAGCCCGGT 1
RESULT 19
AAX24544
ID AAX24544 standard; DNA; 20 BP.
XX
XX AAX24544;
AC
XX
XX 21-JUN-1999 (first entry)
DT
XX
XX Human SR-BI gene exon 8 probe.
DE
XX
XX SR-BI: human; polymorphism; cardiovascular disorder; ischaemia;
XX restenosis; congestive heart failure; atherosclerosis; cholesterol;
XX low density lipoprotein; LDL; high density lipoprotein; HDL;
XX diagnosis; body mass index; obesity; cachexia; gallstone;
XX probe; hybridisation; ss.
XX
XX Synthetic.
XX Homo sapiens.
XX
XX WO9902735-A2.
XX
XX 21-JAN-1999.
XX
XX 10-JUL-1998; 98MO-US14354.
XX
XX 27-FEB-1998; 98US-0031626.
XX
XX 10-JUL-1997; 97US-0890979.
XX
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PA (MILL-) MILLENNIUM PHARM INC.
PA (UYTU-) UNIV TUEFS.
XX
XX Acton SL, Ordovas JM;
PI
XX WPI: 1999-120935/10.
DR
XX
XX
PT Detecting genetic predisposition for body mass disorders - by
PT identifying allelic variants of a polymorphic region of the SR-BI
PT gene
XX
XX Example 2; Page 33; 102pp; English.
PS
XX This probe is designed to detect a C/T polymorphism located at
CC nucleotide 41 of exon 8 of the human SR-BI gene (see AAX24536).
CC It hybridises specifically to the complement of a nucleotide
CC sequence wherein nucleotide 41 is cytidine. The invention is
CC based on the discovery of the genomic structure of the human SR-BI
CC gene (see AAX24498-509) and on the identification of polymorphic
CC regions within the gene which are associated with abnormal body
CC mass index (BMI) and abnormal lipoprotein levels and hence with
CC disorders such as obesity, cachexia, cardiovascular disorders and
CC gallstone formation. The invention provides methods for
CC determining whether a subject has, or is at risk of developing, a
CC disease associated with a specific allele of a polymorphic region
CC of an SR-BI gene. Kits comprising the relevant probe or primer are
CC claimed.
CC
XX Sequence 20 BP; 4 A; 8 C; 4 G; 4 T; 0 other;

Query Match 0.7%; Score 17; DB 20; Length 20;
Best Local Similarity 100.0%; Pred. No. 1.4e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgccgaccggtt 1128
Db 4 tcaacgccgaccggtt 20

RESULT 20
AAX24634/C
ID AAX24634 standard; DNA: 20 BP.
XX
AC AAX24634;
XX
DT 21-JUN-1999 (first entry)
XX
XX Human SR-BI gene exon 8 probe.
DE
XX SR-BI; human; polymorphism; cardiovascular disorder; ischaemia;
KW restenosis; congestive heart failure; atherosclerosis; cholesterol;
KW low density lipoprotein; LDL; high density lipoprotein; HDL;
KW diagnosis; body mass index; obesity; cachexia; gallstone;
KW probe; hybridisation; ss.
XX
XX Synthetic.
OS Homo sapiens.
XX
PN WO9902736-A2.
XX
PD 21-JAN-1999.
XX
PF 10-JUL-1998; 98WO-US14359.
XX
PR 27-FEB-1998; 98US-0032894.
PR 10-JUL-1997; 97US-0890980.
XX
XX (MILL-) MILLENNIUM PHARM INC.
PA Acton SL;
PI
XX
XX WPI: 1999-120936/10.

XX
PT New nucleic acids comprising intronic sequence of a human scavenger
PT receptor-BI (SR-BI) gene - useful for prognosis, diagnosis and
PT treatment of SR-BI associated diseases or conditions
XX
XX Claim 36; Page 32; 103pp; English.
PS
XX

CC This probe is designed to detect a C/T polymorphism located at
CC nucleotide 41 of exon 8 of the human SR-BI gene (see AAX24628).
CC It hybridises specifically to a nucleotide sequence wherein
CC nucleotide 41 of exon 8 is cytidine. The invention is based on
CC the discovery of the genomic structure of the human SR-BI gene (see
CC AAX24590-601) and on the identification of polymorphic regions within
CC the gene which are associated with abnormal body mass index (BMI)
CC and abnormal lipoprotein levels and hence with disorders such as
CC obesity, cachexia, cardiovascular disorders and gallstone formation.
CC The invention provides methods for determining whether a subject
CC has, or is at risk of developing, a disease associated with a
CC specific allele of a polymorphic region of an SR-BI gene. Kits
CC comprising the relevant probe or primer are claimed.
CC
XX

Query Match 0.7%; Score 17; DB 20; Length 20;
Best Local Similarity 100.0%; Pred. No. 1.4e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgccgaccggtt 1128
Db 17 TCAACGCCGACCGGTT 1

RESULT 21
AAX24636
ID AAX24636 standard; DNA: 20 BP.
XX
AC AAX24636;
XX
DT 21-JUN-1999 (first entry)
XX
XX Human SR-BI gene exon 8 probe.
DE
XX SR-BI; human; polymorphism; cardiovascular disorder; ischaemia;
KW restenosis; congestive heart failure; atherosclerosis; cholesterol;
KW low density lipoprotein; LDL; high density lipoprotein; HDL;
KW diagnosis; body mass index; obesity; cachexia; gallstone;
KW probe; hybridisation; ss.
XX
XX Synthetic.
OS Homo sapiens.
XX
PN WO9902736-A2.
XX
PD 21-JAN-1999.
XX
PF 10-JUL-1998; 98WO-US14359.
XX
PR 27-FEB-1998; 98US-0032894.
PR 10-JUL-1997; 97US-0890980.
XX
XX (MILL-) MILLENNIUM PHARM INC.
PA Acton SL;
PI
XX
XX WPI: 1999-120936/10.
DR
XX New nucleic acids comprising intronic sequence of a human scavenger
PT receptor-BI (SR-BI) gene - useful for prognosis, diagnosis and
PT treatment of SR-BI associated diseases or conditions
XX
XX Claim 36; Page 32; 103pp; English.
XX

CC This probe is designed to detect a C/T polymorphism located at
 CC nucleotide 41 of exon 8 of the human SR-BI gene (see AAX24628).
 CC It hybridizes specifically to the complement of a sequence wherein
 CC nucleotide 41 of exon 8 is cytidine. The invention is based on
 CC the discovery of the genomic structure of the human SR-BI gene (see
 CC AAX24590-601) and on the identification of polymorphic regions within
 CC the gene which are associated with abnormal body mass index (BMI)
 CC and abnormal lipoprotein levels and hence with disorders such as
 CC obesity, cachexia, cardiovascular disorders and gallstone formation.
 CC The invention provides methods for determining whether a subject
 CC has, or is at risk of developing, a disease associated with a
 CC specific allele of a polymorphic region of an SR-BI gene. Kits
 CC comprising the relevant probe or primer are claimed.

SO Sequence 20 BP; 4 A; 8 C; 4 G; 4 T; 0 other;

Query Match 0.7%; Score 17; DB 20; Length 20;
 Best Local Similarity 100.0%; Pred. No. 1.4e+03;
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaagccgacccggtt 1128
 |||||
 DB 4 tcaagccgacccggtt 20

RESULT 22
 AAQ75787/c
 ID AAQ75787 standard; DNA; 21 BP.
 XX
 AC AAQ75787;
 XX
 DT 04-AUG-1995 (first entry)

DE Reverse transcription primer used in cDNA analysis technique.
 XX
 XX Analysis: gene expression; reverse transcription; primer; cDNA;
 KW aggregate; restriction enzyme; ss.
 XX
 OS Synthetic.

PN JP06303997-A.
 XX
 PD 01-NOV-1994.
 XX
 PF 16-APR-1993; 93JP-0112515.
 XX
 PR 16-APR-1993; 93JP-0112515.
 XX

PA (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.
 XX
 DR WPI; 1995-018287/03.
 XX

PT Analysis of cDNA and gene expression - by amplification of mRNA
 PT followed by digestion with restriction enzymes
 XX
 PS Disclosure; Page 9; 11pp; Japanese.

XX A method for the analysis of cDNA comprises (a) preparing an
 CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs
 CC and a plural type of labelled reverse transcription primers
 CC (GENESEQ files AAQ75547-075798) and using the aggregate of mRNAs as the
 CC template for each reverse transcription primer; (b) digesting each of
 CC the prepared aggregates of the double-stranded cDNAs with restriction
 CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
 CC separate lanes. The method can be used to analyse gene expression
 CC rapidly and easily.

SO Sequence 21 BP; 1 A; 2 C; 1 G; 17 T; 0 other;

Query Match 0.7%; Score 17; DB 16; Length 21;
 Best Local Similarity 100.0%; Pred. No. 1.4e+03;

Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2550 tggaaaaaaaaaaaaa 2566
 |||||
 DB 20 TGGAAAAAAAAAAAAA 4

RESULT 23
 AAQ75788/c
 ID AAQ75788 standard; DNA; 21 BP.
 XX
 AC AAQ75788;
 XX

DT 04-AUG-1995 (first entry)

DE Reverse transcription primer used in cDNA analysis technique.

XX Analysis: gene expression; reverse transcription; primer; cDNA;
 KW aggregate; restriction enzyme; ss.

OS Synthetic.

PN JP06303997-A.
 XX
 PD 01-NOV-1994.
 XX

PF 16-APR-1993; 93JP-0112515.
 XX
 PR 16-APR-1993; 93JP-0112515;
 XX

PA (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.
 XX

DR WPI; 1995-018287/03.

PT Analysis of cDNA and gene expression - by amplification of mRNA
 PT followed by digestion with restriction enzymes
 XX
 PS Disclosure; Page 9; 11pp; Japanese.

XX A method for the analysis of cDNA comprises (a) preparing an
 CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs
 CC and a plural type of labelled reverse transcription primers
 CC (GENESEQ files AAQ75547-075798) and using the aggregate of mRNAs as the
 CC template for each reverse transcription primer; (b) digesting each of
 CC the prepared aggregates of the double-stranded cDNAs with restriction
 CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
 CC separate lanes. The method can be used to analyse gene expression
 CC rapidly and easily.

SO Sequence 21 BP; 2 A; 2 C; 0 G; 17 T; 0 other;

Query Match 0.7%; Score 17; DB 16; Length 21;
 Best Local Similarity 100.0%; Pred. No. 1.4e+03;
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2550 tggaaaaaaaaaaaaa 2566
 |||||
 DB 20 TGGAAAAAAAAAAAAA 4

RESULT 24
 AAQ75790/c
 ID AAQ75790 standard; DNA; 21 BP.
 XX
 AC AAQ75790;
 XX

DT 04-AUG-1995 (first entry)

DE Reverse transcription primer used in cDNA analysis technique.

XX Analysis: gene expression; reverse transcription; primer; cDNA;
 KW aggregate; restriction enzyme; ss.

XX	RESULT
XX	25
XX	AAC96163/C
ID	AAC96163 standard; DNA; 25 BP.
XX	
XX	AAC96163;
AC	
XX	
DT	26-FEB-2001 (first entry)
XX	
DE	16s rRNA gene PCR primer #130.
XX	
XX	DNA sequence analysis; sequencing; protein sequence; protein structure;
KW	gene typing; organ donation; bacteria identification; 16s rRNA; HLA;
KV	human leukocyte antigen; PCR primer; ss.
XX	
OS	Homo sapiens.
XX	
PN	WO200065088-A2.
XX	
PD	02-NOV-2000.
XX	
PF	20-APR-2000; 2000WO-EP03636.
XX	
PR	26-APR-1999; 99EP-0303215.
XX	
PA	(AMSH) AMERSHAM PHARMACIA BIOTECH AB.
XX	
PI	Ulfendahl P, Wong K;
XX	
DR	WPI: 2000-679677/66.
XX	
DT	Identifying extendible primers for use in identification, or

CC Oligonucleotides characterized by the method form pharmaceutical

CC proliferation, and being active against a eukaryotic pathogen, a human
 CC retrovirus, a human immunodeficiency virus (HIV), or a non-human
 CC retrovirus, including influenza virus, Epstein-Barr virus, Respiratory
 CC Syncytial Virus or cytomegalovirus (CMV). The compositions enable
 CC characterization of deletion sequence oligonucleotides having related
 CC but different nucleobase sequences, and quantification of different
 CC species of deletion sequence ("target") oligonucleotides in a mixture.
 CC Also, if the specificity of the oligonucleotide's nucleobase sequence
 CC for its reverse complement is not modified, the method may be performed
 CC using oligodeoxynucleotides.

SQ Sequence 27 BP; 7 A; 2 C; 3 G; 15 T; 0 other;

Query Match 0.7%; Score 17; DB 20; Length 27;
 Best Local Similarity 100.0%; Pred. No. 1.4e+03;
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2550 tgaataaaataaaataaa 2566
 ||||||||||||||||
 DB 18 TGAATAAAATAAAATAAA 2

RESULT 27
 AAF86334
 ID AAF86334 standard; DNA; 31 BP.

AC AAF86334;
 DT 17-JUL-2001 (first entry)
 DE PCR primer 3 specific for human ZFP-52 cDNA.

KW Human; zinc finger protein 52; ZFP-52; proliferation disorder; neoplasm;
 KW immune system disorder; metabolic disorder; cancer; PCR primer; ss.

OS Homo sapiens.

PN W0200127151-A1.

PD 19-APR-2001.

PF 08-OCT-2000; 2000WO-CN00308.

PR 10-OCT-1999; 99CN-0116949.

PA (SHAN-) SHANGHAI BIO DOOR GENE TECHNOLOGY LTD.

PI Mao Y, Xie Y;

DR WPI; 2001-281977/29.

PT Human zinc finger protein 52 applicable in diagnosis and treatment of
 PT proliferation disorders, disorders induced by immune system, metabolic
 PT disorders, neoplasms and cancers

PS Example 5; Page 14; 30pp; Chinese.

CC This invention relates to human zinc finger protein 52 (ZFP-52) which is
 CC a member of the kruppel family of proteins. This sequence represents a
 CC PCR primer used to amplify cDNA encoding human ZFP-52. ZFP-52 protein and
 CC polynucleotide sequences can be used in the diagnosis and treatment of
 CC proliferation disorders, immune system related disorders, metabolic
 CC disorders, neoplasms and cancer.

SQ Sequence 31 BP; 16 A; 5 C; 6 G; 4 T; 0 other;

Query Match 0.7%; Score 17; DB 22; Length 31;
 Best Local Similarity 100.0%; Pred. No. 1.3e+03;
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2549 atggaataaaataaaataaa 2565

DB 10 atggaataaaataaaataaa 26
 ||||||||||||||||

RESULT 28
 AAX28398/c
 ID AAX28398 standard; DNA; 47 BP.

AC AAX28398;

DT 21-JUN-1999 (first entry)

DE Primer for CCR5 gene.

KW Primer; CCR5 gene; non-synctia-inducing; HIV-1; mutation detection;
 KW chemokine receptor gene; infection; disease progression prediction; ss.

OS Synthetic.

PN W09913112-A1.

PD 18-MAR-1999.

PF 14-SEP-1998; 98WO-US19007.

PR 12-SEP-1997; 97US-0928465.

PA (ALKU) AKZO NOBEL NV.

PI Lee EM, Romano JW;

DR WPI; 1999-263372/22.

PT Determination of zygosity of CCR5 chemokine receptor gene in an
 PT individual

PS Claim 6; Page 23; 36pp; English.

CC This sequence represents a primer for a region of the CCR5 gene.
 CC The invention relates to a method for the determination of susceptibility
 CC of an individual to non-synctia-inducing (NSI) forms of human
 CC immunodeficiency virus type 1 (HIV-1), by detecting whether the
 CC individual is homozygous mutant, heterozygous or homozygous wild type for
 CC the CCR5 chemokine receptor gene. The method can be used to predict
 CC susceptibility of an individual to infection by NSI forms of HIV-1 and
 CC for predicting disease progression.

SQ Sequence 47 BP; 15 A; 14 C; 11 G; 7 T; 0 other;

Query Match 0.7%; Score 17; DB 20; Length 47;
 Best Local Similarity 100.0%; Pred. No. 1.3e+03;
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1295 tgcctgcgcgcgcgc 1311
 ||||||||||||||||
 DB 41 TGGTCTGCGCGCTGCTC 25

RESULT 29
 AAX69802/c
 ID AAX69802 standard; RNA; 17 BP.

AC AAX69802;

DT 28-JUL-1999 (first entry)

DE Human flt1 VEGF receptor hammerhead ribozyme substrate #1097.

KW Vascular endothelial growth factor receptor; VEGF receptor; flt-1;
 KW flt-1; KDR; hammerhead ribozyme; hairpin ribozyme; cleavage;
 KW tumour angiogenesis; psoriasis; rheumatoid arthritis; ocular disease;
 KW fms-like tyrosine kinase 1; kinase insert domain containing receptor;

XX	PR	11-JAN-1996;	96US-0584040.
XX	PR	26-OCT-1995;	95US-0005974.
XX	PA	(CHIR) CHIRON CORP.	
XX	PI	(RIBO-) RIBOSYME PHARM INC.	
XX	PJ	Escobedo J, McSwiggen J, Pavco P, Slinchcomb D;	
XX	DR	WPI: 1997-259017/23.	
XX	PT	Nucleic acid molecule modulating VEGF receptor(s) gene expression or	
XX	PT	mRNA stability - useful for treating e.g. tumour angiogenesis,	
XX	PT	psoriasis, rheumatoid arthritis, etc., in a human patient	
XX	PS	Claim 4; Page 79; 218pp; English.	
XX	CC	The present invention describes nucleic acid molecules which modulate	
XX	CC	the synthesis, expression and/or stability of a mRNA encoding 1 or more	
XX	CC	receptors of vascular endothelial growth factor (VEGF). A patient	
XX	CC	(preferably human) having a condition associated with the level of the	
XX	CC	fms-like tyrosine kinase 1 (flt-1), kinase insert domain containing	
XX	CC	receptor (KDR) and/or foetal liver kinase 1 (flk-1) (e.g. tumour	
XX	CC	angiogenesis, ocular diseases, psoriasis and rheumatoid arthritis) can	
XX	CC	be treated by administering the nucleic acid molecule or the expression	
XX	CC	vector to the patient. AA67275 to AA75752 represent specific examples	
XX	CC	of nucleic acid molecules from the present invention.	
SQ		Sequence 17 BP; 2 A; 2 C; 0 G; 13 U; 0 other;	
OY	Query Match	0.6%; Score 16; DB 18; Length 17;	
	Best Local Similarity	100.0%; Pred. No. 3.0e+03;	
	Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;		
DB	2550 tggaaaaaaataaaa 2565		
	16 TGGAAAAAAAAAAAAA 1		
RESULT 31			
ID	AAV54175/C		
	AAV54175 standard; CDNA: 18 BP.		
XX	AC	AAV54175;	
XX	DT	21-DEC-1998 (first entry)	
XX	DE	Nucleotide sequence PCR primer 12.	
XX	KW	PCR, primer: amplification; apoptosis; antibody; inhibition; ss;	
XX	KM	immunohistological staining.	
XX	OS	Synthetic.	
XX	PN	WO9839437-A1.	
XX	PD	11-SEP-1998.	
XX	PF	05-MAR-1998; 98WO-JP00905.	
XX	PR	05-MAR-1997; 97JP-0050302.	
XX	PA	(KYOW) KYOWA HAKKO KOGYO KK.	
XX	PI	Sakaki Y;	
XX	DR	WPI: 1998-495844/42.	
XX	PT	Novel apoptosis-related DNAs and proteins for diagnosis,	
XX	PT	preventing or treating diseases associated with apoptosis	
XX	PS	Example 1; Page 51; 70pp; Japanese.	
XX			

CC This is the nucleotide sequence of a PCR primer used in the method
CC of the invention, involving the use of novel apoptosis-related DNAs
CC and proteins. The inventions can be used as diagnostic reagents for
CC apoptosis e.g. (monoclonal) antibodies for the protein, as a reagent
CC in immunohistological staining, as apoptosis inhibitors. It can also
CC be used for treatment of apoptosis-related diseases.
XX
SO

Sequence 18 BP; 0 A; 2 C; 1 G; 15 T; 0 other;

Query Match 0.6%; Score 16; DB 19; Length 18;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaataaaataaa 2566
DB 18 GGAATAAATAAATAAATAA 3

RESULT 32

AAA58385
ID AAA58385 standard; DNA; 18 BP.

AC AAA58385;

DT 01-NOV-2000 (first entry)

DE Polynucleotide # 1 used in a biomolecule detection system.

KW Nanocrystal; biomolecule detection; nonisotopic detection system; ss.

OS Synthetic.

PN W0200028088-A1.

PD 18-MAY-2000.

PF 10-NOV-1999; 99WO-US26612.

PR 10-NOV-1998; 98US-0107828.

PR 09-NOV-1999; 99US-0437076.

PA (BIOC-) BIOCRYSTAL LTD.

PI Barbera-Guillem E, Nelson MB, Castro S;

DR WPI; 2000-376593/32.

PT Functionalized nanocrystal carrying polynucleotide, used for detecting
PT target analyte, forms dendrimers with complementary nanocrystals to
PT amplify the fluorescent signal

PS Example 3; Page 68; 72pp; English.

CC The present invention relates to functionalised nanocrystals for use in
CC nonisotopic detection systems for biomolecules e.g. nucleic acids,
CC proteins, lipids or drugs. The nanocrystals have polynucleotide strands
CC attached to their surfaces with one end of the polynucleotide extending
CC outwardly from the nanocrystal. The present sequence is one such
CC polynucleotide. These nanocrystals are used with a second series of
CC nanocrystals, which have polynucleotides complementary to the first
CC polynucleotides, so that the respective complementary strands hybridise
CC to each other and form a dendrimer. This dendrimer produces a signal
CC which can then be detected e.g. fluorescence. The present sequence is
CC composed mainly of adenine bases. This sequence may therefore be
CC used with a polynucleotide composed mainly of thymine bases (AAA58386).
XX
SO

Sequence 18 BP; 15 A; 0 C; 3 G; 0 T; 0 other;

Query Match 0.6%; Score 16; DB 21; Length 18;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaataaaataaa 2566

DB 2 ggaataaaataaaataaa 17

RESULT 33

AAZ90651/C
ID AAZ90651 standard; DNA; 18 BP.

AC AAZ90651;

DT 13-JUN-2000 (first entry)

DE Human adipose tissue gene amplifying primer #12.

KW Adipose tissue; obesity; diabetes; hyperlipemia; hypertension; human;
KW arteriosclerosis; hyperuricemia; sleep apnea syndrome; PCR primer; ss.

OS Homo sapiens.

PN JP2000037190-A.

PD 08-FEB-2000.

PF 23-JUL-1998; 98JP-0225228.

PR 23-JUL-1998; 98JP-0225228.

PA (NISH) JAPAN TOBACCO INC.

DR WPI; 2000-306578/27.

PT A physiologically active protein specifically derived from mammal
PT tissue

PS Example 2; Page 18; 50pp; Japanese.

CC The invention relates to identification of genes and proteins of adipose
CC tissue relating to obesity, particularly complications of visceral
CC obesity including diabetes, hyperlipemia, hypertension,
CC arteriosclerosis, hyperuricemia and sleep apnea syndrome. The genes
CC (AAZ90631-633) and the proteins (AAZ90631-633) are used in the genetic
CC diagnosis, prevention and treatment of adipose tissue related diseases.
CC Sequences AAZ90640-51 represent PCR primers amplifying the human adipose
CC tissue genes.
XX
SO

Sequence 18 BP; 0 A; 2 C; 1 G; 15 T; 0 other;

Query Match 0.6%; Score 16; DB 21; Length 18;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaataaaataaa 2566
DB 18 GGAATAAATAAATAAATAA 3

RESULT 34

AAQ75558/C
ID AAQ75558 standard; DNA; 19 BP.

AC AAQ75558;

DT 04-AUG-1995 (first entry)

DE Reverse transcription primer used in cDNA analysis technique.

XX Analysis; gene expression; reverse transcription; primer; cDNA;
KW aggregate; restriction enzyme; ss.
XX
OS Synthetic.

```
XX JP06303997-A.
PN 01-NOV-1994.
XX 16-APR-1993; 93JP-0112515.
XX 16-APR-1993; 93JP-0112515.
PR 16-APR-1993; 93JP-0112515.
XX (NITE ) NIPPON TELEGRAPH & TELEPHONE CORP.
XX WPI; 1995-018287/03.
XX Analysis of cDNA and gene expression - by amplification of mRNA
PT followed by digestion with restriction enzymes
XX Disclosure; Page 5; 11pp; Japanese.
XX A method for the analysis of cDNA comprises (a) preparing an
CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs
CC and a plural type of labelled reverse transcription primers
CC (GENESEQ files AAQ75547-075798) and using the aggregate of mRNAs as the
CC template for each reverse transcription primer; (b) digesting each of
CC the prepared aggregates of the double-stranded cDNAs with restriction
CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
CC separate lanes. The method can be used to analyse gene expression
CC rapidly and easily.
XX Sequence 19 BP; 0 A; 2 C; 0 G; 17 T; 0 other;
SQ
Query Match 0.6%; Score 16; DB 16; Length 19;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
OY 2551 ggaataaaaaaa 2566
DB 19 GGAATAAAAAAAAA 4
RESULT 35
AAQ75603/c
ID AAQ75603 standard; DNA; 20 BP.
XX AAQ75603;
AC AAQ75603;
XX 04-AUG-1995 (first entry)
DT 04-AUG-1995 (first entry)
XX Reverse transcription primer used in cDNA analysis technique.
DE Reverse transcription primer used in cDNA analysis technique.
XX Analysis; gene expression; reverse transcription; primer; cDNA;
KM aggregate; restriction enzyme; ss.
XX Synthetic.
OS Synthetic.
XX JP06303997-A.
PN 01-NOV-1994.
XX 16-APR-1993; 93JP-0112515.
XX 16-APR-1993; 93JP-0112515.
PR 16-APR-1993; 93JP-0112515.
XX (NITE ) NIPPON TELEGRAPH & TELEPHONE CORP.
XX WPI; 1995-018287/03.
XX Analysis of cDNA and gene expression - by amplification of mRNA
PT followed by digestion with restriction enzymes
XX Disclosure; Page 5; 11pp; Japanese.
XX A method for the analysis of cDNA comprises (a) preparing an
```

```
CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs
CC and a plural type of labelled reverse transcription primers
CC (GENESEQ files AAQ75547-075798) and using the aggregate of mRNAs as the
CC template for each reverse transcription primer; (b) digesting each of
CC the prepared aggregates of the double-stranded cDNAs with restriction
CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
CC separate lanes. The method can be used to analyse gene expression
CC rapidly and easily.
XX Sequence 20 BP; 0 A; 2 C; 1 G; 17 T; 0 other;
SQ
Query Match 0.6%; Score 16; DB 16; Length 20;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
OY 2551 ggaataaaaaaa 2566
DB 19 GGAATAAAAAAAAA 4
RESULT 36
AAQ75605/c
ID AAQ75605 standard; DNA; 20 BP.
XX AAQ75605;
AC AAQ75605;
XX 04-AUG-1995 (first entry)
DT 04-AUG-1995 (first entry)
XX Reverse transcription primer used in cDNA analysis technique.
DE Reverse transcription primer used in cDNA analysis technique.
XX Analysis; gene expression; reverse transcription; primer; cDNA;
KM aggregate; restriction enzyme; ss.
XX Synthetic.
OS Synthetic.
XX JP06303997-A.
PN 01-NOV-1994.
XX 16-APR-1993; 93JP-0112515.
XX 16-APR-1993; 93JP-0112515.
PR 16-APR-1993; 93JP-0112515.
XX (NITE ) NIPPON TELEGRAPH & TELEPHONE CORP.
XX WPI; 1995-018287/03.
XX Analysis of cDNA and gene expression - by amplification of mRNA
PT followed by digestion with restriction enzymes
XX Disclosure; Page 5; 11pp; Japanese.
XX A method for the analysis of cDNA comprises (a) preparing an
CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs
CC and a plural type of labelled reverse transcription primers
CC (GENESEQ files AAQ75547-075798) and using the aggregate of mRNAs as the
CC template for each reverse transcription primer; (b) digesting each of
CC the prepared aggregates of the double-stranded cDNAs with restriction
CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
CC separate lanes. The method can be used to analyse gene expression
CC rapidly and easily.
XX Sequence 20 BP; 0 A; 2 C; 0 G; 18 T; 0 other;
SQ
Query Match 0.6%; Score 16; DB 16; Length 20;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
OY 2551 ggaataaaaaaa 2566
DB 19 GGAATAAAAAAAAA 4
```


RESULT 37
AA075606/c
ID AA075606 standard; DNA: 20 BP.
XX
XX
XX
AC AA075606:
XX
XX
DT 04-AUG-1995 (first entry)
XX
DE Reverse transcription primer used in cDNA analysis technique.
XX
XX Analysis; gene expression; reverse transcription; primer; cDNA;
KW aggregate; restriction enzyme; ss.
XX
XX
OS Synthetic.
XX
PN JP06303997-A.
XX
XX
PD 01-NOV-1994.
XX
XX
PE 16-APR-1993: 93JP-0112515.
XX
XX
PR 16-APR-1993: 93JP-0112515.
XX
XX
PA (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.
XX
XX
DR WPI: 1995-018287/03.
XX
XX Analysis of cDNA and gene expression - by amplification of mRNA
PT followed by digestion with restriction enzymes.
XX
XX
PS Disclosure; Page 5, 11pp; Japanese.
XX
XX
CC A method for the analysis of cDNA comprises (a) preparing an
CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs
CC and a plural type of labelled reverse transcription primers
CC (GENSEQ files AA075547-075798) and using the aggregate of mRNAs as the
CC template for each reverse transcription primer; (b) digesting each of
CC the prepared aggregates of the double-stranded cDNAs with restriction
CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
CC separate lanes. The method can be used to analyse gene expression
CC rapidly and easily.
XX
XX
SQ Sequence 20 BP; 0 A; 3 C; 0 G; 17 T; 0 other;

Query Match 0.6%; Score 16; DB 16; Length 20;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 2551 ggaataaaaaa 2566
DB 19 GGAATAAAAAA 4

RESULT 38
AAZ31280/c
ID AAZ31280 standard; DNA: 20 BP.
XX
XX
AC AAZ31280:
XX
XX
DT 24-JAN-2000 (first entry)
XX
XX
DE CCR5 gene inhibiting antisense oligo AS(s)-37.
XX
XX HIV cofactor inhibitor; HIV infection; CXCR4 gene; CCR5 gene;
KW drug composition; antisense; ss.
XX
XX
OS Synthetic.
XX
XX
PN WO951751-A1.

PD 14-OCT-1999.
XX
XX
PE 01-APR-1999: 99WO-JP01722.
XX
XX
PR 02-APR-1998: 98JP-0125452.
XX
XX
PA (MARI-) MARINE BIO CO LTD.
XX
XX
PI Takaku H, Yamamoto N, Kimura T, Takai K, Wada A;
DR WPI: 1999-620207/53.
XX
XX
PT Antisense oligonucleotide-based HIV cofactor inhibitors, as drug
PT compositions for treatment of HIV infection
XX
XX
PS Claim 6; Page 16; 59pp; Japanese.
XX
XX
CC The invention provides HIV cofactor inhibitors that contain
CC oligonucleotides with a base sequence complementary to the CXCR4 or CCR5
CC genes. Such inhibitors can be formulated into drug compositions for
CC prevention or treatment of HIV infection, with inhibition of expression
CC of CXCR4 or/and CCR5 gene. Sequences AAZ31244-306 represent antisense
CC oligonucleotides to the CCR5 gene.
XX
XX
SQ Sequence 20 BP; 5 A; 8 C; 7 G; 0 U; 0 other;

Query Match 0.6%; Score 16; DB 20; Length 20;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1295 tggctcgcgcgtct 1310
DB 16 TGGCTCGCGCTCT 1

RESULT 39
AAZ72142/c
ID AAZ72142 standard; DNA: 20 BP.
XX
XX
AC AAZ72142:
XX
XX
DT 10-SEP-2001 (first entry)
XX
XX
DE human biallelic marker upstream amplification primer SEQ ID NO:6498.
XX
XX
KW Human genome; biallelic marker; high density disequilibrium map;
KW genomic map; haplotype; phenotype; polymorphic base; genotyping;
KW haplotyping; hybridisation; identification; characterisation;
KW amplification; single nucleotide polymorphism; SNP; PCR primer;
KW diagnosis; ss.
XX
XX
OS Homo sapiens.
XX
XX
PN WO954500-A2.
XX
XX
PD 28-OCT-1999.
XX
XX
PE 21-APR-1999: 99WO-IB00822.
XX
XX
PR 21-APR-1998: 98US-0082614.
PR 23-NOV-1998: 98US-0109732.
XX
XX
PA (GEST) GENSET.
XX
XX
PI Cohen D, Blumenfeld M, Chumakov I;
DR WPI: 2000-013267/01.
XX
XX
PT Novel biallelic markers used to construct a high density disequilibrium
PT map of the human genome
XX
XX
PS Claim 9; Page 1616; 2745pp; English.

XX AA265654 to AA269578 represent human biallelic markers from the present
CC invention, which contain a polymorphic base at position 24 of their
CC nucleotide sequences. AA269579 to AA277440 represent amplification
CC primers for the biallelic markers. The biallelic markers of the
CC invention have a variety of uses: they can be used for high density
CC mapping of the human genome, and in complex association studies and
CC haplotyping studies which are useful in determining the genetic basis
CC for disease states. Compositions and methods of the invention can also
CC be useful for the identification of the targets for the development of
CC pharmaceutical agents and diagnostic methods, as well as the
CC characterization of the differential efficacious responses to and side
CC effects from pharmaceutical agents acting on a disease as well as other
CC treatment.
CC N.B. The SEQ ID NOS 2852, 2913, 2974, 3035, 3096, 3157, 3227, 3297
CC and 3367, are not actually given a sequence in the Sequence Listing
CC from the present invention.
CC
CC
SQ Sequence 20 BP; 4 A; 3 C; 8 G; 5 T; 0 other;

Query Match 0.6%; Score 16; DB 21; Length 20;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1710 caccagctgagctcc 1725
|||||
DB 20 CACAGCCTGAGCCTCC 5

RESULT 40
AAQ75791/c
ID AAQ75791 standard; DNA: 21 BP.
AC AAQ75791;
XX
DT 04-AUG-1995 (first entry)
XX
DE Reverse transcription primer used in cDNA analysis technique.
XX
XX Analysis: gene expression; reverse transcription; primer: cDNA;
KW aggregate; restriction enzyme; ss.
XX
OS Synthetic.
XX
PN JP06303997-A.
XX
PD 01-NOV-1994.
XX
PF 16-APR-1993; 93JP-0112515.
XX
PR 16-APR-1993; 93JP-0112515.
XX
PA (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.
XX
DR WPI; 1995-018287/03.
XX
PT Analysis of cDNA and gene expression - by amplification of mRNA
PT followed by digestion with restriction enzymes
XX
PS Disclosure; Page 9; 11pp; Japanese.
XX
XX A method for the analysis of cDNA comprises (a) preparing an
CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs
CC and a plural type of labelled reverse transcription primers
CC (GENESEQ files AAQ7547-075798) and using the aggregate of mRNAs as the
CC template for each reverse transcription primer; (b) digesting each of
CC the prepared aggregates of the double-stranded cDNAs with restriction
CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
CC separate lanes. The method can be used to analyse gene expression
CC rapidly and easily.
XX
XX Sequence 21 BP; 0 A; 2 C; 1 G; 18 T; 0 other;

Query Match 0.6%; Score 16; DB 16; Length 21;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaaaa 2566
|||||
DB 19 GGAATAAAAAAAAAA 4

RESULT 41
AAQ75792/c
ID AAQ75792 standard; DNA: 21 BP.
AC AAQ75792;
XX
DT 04-AUG-1995 (first entry)
XX
DE Reverse transcription primer used in cDNA analysis technique.
XX
XX Analysis: gene expression; reverse transcription; primer: cDNA;
KW aggregate; restriction enzyme; ss.
XX
OS Synthetic.
XX
PN JP06303997-A.
XX
PD 01-NOV-1994.
XX
PF 16-APR-1993; 93JP-0112515.
XX
PR 16-APR-1993; 93JP-0112515.
XX
PA (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.
XX
DR WPI; 1995-018287/03.
XX
PT Analysis of cDNA and gene expression - by amplification of mRNA
PT followed by digestion with restriction enzymes
XX
PS Disclosure; Page 9; 11pp; Japanese.
XX
XX A method for the analysis of cDNA comprises (a) preparing an
CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs
CC and a plural type of labelled reverse transcription primers
CC (GENESEQ files AAQ7547-075798) and using the aggregate of mRNAs as the
CC template for each reverse transcription primer; (b) digesting each of
CC the prepared aggregates of the double-stranded cDNAs with restriction
CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
CC separate lanes. The method can be used to analyse gene expression
CC rapidly and easily.
XX
XX Sequence 21 BP; 1 A; 2 C; 0 G; 18 T; 0 other;

Query Match 0.6%; Score 16; DB 16; Length 21;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaaaa 2566
|||||
DB 19 GGAATAAAAAAAAAA 4

RESULT 42
AAQ75793/c
ID AAQ75793 standard; DNA: 21 BP.
AC AAQ75793;
XX
DT 04-AUG-1995 (first entry)
XX

DE Reverse transcription primer used in cDNA analysis technique.
 XX
 KW Analysis; gene expression; reverse transcription; primer; cDNA;
 KM aggregate; restriction enzyme; ss.
 XX
 OS Synthetic.
 XX
 PN JP06303997-A.
 XX
 PD 01-NOV-1994.
 XX
 PF 16-APR-1993; 93JP-0112515.
 XX
 PR 16-APR-1993; 93JP-0112515.
 XX
 PA (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.
 XX
 DR WPI; 1995-018287/03.
 XX
 PT Analysis of cDNA and gene expression - by amplification of mRNA
 PT followed by digestion with restriction enzymes
 XX
 PS Disclosure; Page 9; 11pp; Japanese.
 XX
 CC A method for the analysis of cDNA comprises (a) preparing an
 CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs
 CC and a plural type of labelled reverse transcription primers
 CC (GENESKO files AAQ75547-Q75798) and using the aggregate of mRNAs as the
 CC template for each reverse transcription primer; (b) digesting each of
 CC the prepared aggregates of the double-stranded cDNAs with restriction
 CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
 CC separate lanes. The method can be used to analyse gene expression
 CC rapidly and easily.
 XX
 SQ Sequence 21 BP; 0 A; 2 C; 0 G; 19 T; 0 other;
 XX

Query Match 0.6%; Score 16; DB 16; Length 21;
 Best Local Similarity 100.0%; Pred. No. 3.7e+03;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaataaaataaa 2566
 ||||||||||||||||
 Db 19 GGAATAAAATAAAATAAA 4

RESULT 43
 AAQ75794/c
 ID AAQ75794 standard; DNA: 21 BP.
 XX
 AC AAQ75794;
 XX
 DT 04-AUG-1995 (first entry)
 XX
 DE Reverse transcription primer used in cDNA analysis technique.
 XX
 KM Analysis; gene expression; reverse transcription; primer; cDNA;
 KM aggregate; restriction enzyme; ss.
 XX
 OS Synthetic.
 XX
 PN JP06303997-A.
 XX
 PD 01-NOV-1994.
 XX
 PF 16-APR-1993; 93JP-0112515.
 XX
 PR 16-APR-1993; 93JP-0112515.
 XX
 PA (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.
 XX
 DR WPI; 1995-018287/03.
 XX

PT Analysis of cDNA and gene expression - by amplification of mRNA
 PT followed by digestion with restriction enzymes
 XX
 PS Disclosure; Page 9; 11pp; Japanese.
 XX
 CC A method for the analysis of cDNA comprises (a) preparing an
 CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs
 CC and a plural type of labelled reverse transcription primers
 CC (GENESKO files AAQ75547-Q75798) and using the aggregate of mRNAs as the
 CC template for each reverse transcription primer; (b) digesting each of
 CC the prepared aggregates of the double-stranded cDNAs with restriction
 CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in
 CC separate lanes. The method can be used to analyse gene expression
 CC rapidly and easily.
 XX
 SQ Sequence 21 BP; 0 A; 3 C; 0 G; 18 T; 0 other;
 XX

Query Match 0.6%; Score 16; DB 16; Length 21;
 Best Local Similarity 100.0%; Pred. No. 3.7e+03;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaataaaataaa 2566
 ||||||||||||||||
 Db 19 GGAATAAAATAAAATAAA 4

RESULT 44
 AAQ75795/c
 ID AAQ75795 standard; DNA: 21 BP.
 XX
 AC AAQ75795;
 XX
 DT 04-AUG-1995 (first entry)
 XX
 DE Reverse transcription primer used in cDNA analysis technique.
 XX
 KM Analysis; gene expression; reverse transcription; primer; cDNA;
 KM aggregate; restriction enzyme; ss.
 XX
 OS Synthetic.
 XX
 PN JP06303997-A.
 XX
 PD 01-NOV-1994.
 XX
 PF 16-APR-1993; 93JP-0112515.
 XX
 PR 16-APR-1993; 93JP-0112515.
 XX
 PA (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.
 XX
 DR WPI; 1995-018287/03.
 XX

Query Match 0.6%; Score 16; DB 16; Length 21;
 Best Local Similarity 100.0%; Pred. No. 3.7e+03;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaataaaataaa 2566
 ||||||||||||||||
 Db 19 GGAATAAAATAAAATAAA 4

RESULT 44
 AAQ75795/c
 ID AAQ75795 standard; DNA: 21 BP.
 XX
 AC AAQ75795;
 XX
 DT 04-AUG-1995 (first entry)
 XX
 DE Reverse transcription primer used in cDNA analysis technique.
 XX
 KM Analysis; gene expression; reverse transcription; primer; cDNA;
 KM aggregate; restriction enzyme; ss.
 XX
 OS Synthetic.
 XX
 PN JP06303997-A.
 XX
 PD 01-NOV-1994.
 XX
 PF 16-APR-1993; 93JP-0112515.
 XX
 PR 16-APR-1993; 93JP-0112515.
 XX
 PA (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.
 XX
 DR WPI; 1995-018287/03.
 XX

Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaaaaaa.2566
|||||
Db 19 GGAATAAAAAAAAA 4

RESULT 45

AA075796/c
ID AA075796 standard; DNA: 21 BP.

XX AA075796;

DT 04-AUG-1995 (first entry)

DE Reverse transcription primer used in cDNA analysis technique.

XX Analysis; gene expression; reverse transcription; primer; cDNA;

KM aggregate; restriction enzyme; ss.

XX Synthetic.

OS JP06303997-A.

XX 01-NOV-1994.

PD 16-APR-1993; 93JP-0112515.

XX 16-APR-1993; 93JP-0112515.

XX (NITE) NIPPON TELEGRAPH & TELEPHONE CORP.

DR WPI; 1995-018287/03.

XX Analysis of cDNA and gene expression - by amplification of mRNA

PT followed by digestion with restriction enzymes

XX Disclosure; Page 9; 11pp; Japanese.

XX A method for the analysis of cDNA comprises (a) preparing an

CC aggregate of double-stranded cDNAs by using an aggregate of mRNAs

CC and a plural type of labelled reverse transcription primers

CC (GENESIO files AA075547-075798) and using the aggregate of mRNAs as the

CC template for each reverse transcription primer; (b) digesting each of

CC the prepared aggregates of the double-stranded cDNAs with restriction

CC enzyme and; (c) electrophoresing the digested aggregate of cDNAs in

CC separate lanes. The method can be used to analyse gene expression

CC rapidly and easily.

XX Sequence 21 BP: 1 A; 3 C; 0 G; 17 T; 0 other;

Query Match 0.6%; Score 16; DB 16; Length 21;
Best Local Similarity 100.0%; Pred. No. 3.7e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaaaaaa.2566
|||||
Db 19 GGAATAAAAAAAAA 4

Search completed: April 20, 2002, 10:14:07
Job time: 6208 sec

GenCore version 4.5
Copyright (c) 1993 - 2000 CompuGen Ltd.

OM nucleic - nucleic search, using sw model

Run on: April 20, 2002, 06:42:18 ; Search time 2150.98 Seconds
(without alignments)
12819.119 Million cell updates/sec

Title: US-10-024-396-3

Perfect score: 2566

Sequence: 1 cytcgcgcgtccctctctcct.....aaatgaaaaaaaaaaaaa 2566

Scoring table:

OLIGO_NUC
Gapop 60.0, Gapext 60.0

Searched: 11351937 seqs, 537289281 residues

Word size: 0

Total number of hits satisfying chosen parameters: 80718

Minimum DB seq length: 0
Maximum DB seq length: 50

Post-processing: Listing first 45 summaries

Database:

EST:
1: em_estfun:
2: em_esthum:
3: em_estlin:
4: em_estom:
5: em_estopl:
6: em_estlba:
7: em_estlro:
8: em_estov:
9: em_hic:
10: qb_estl:
11: qb_est2:
12: qb_hic:
13: qb_gss:
14: em_gss_fun:
15: em_gss_hum:
16: em_gss_lin:
17: em_gss_pln:
18: em_gss_pro:
19: em_gss_pro:
20: em_gss_vit:
21: em_gss_other:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	34	1.3	40	11	R71941
2	19	0.7	19	13	A2662226
3	19	0.7	48	13	A2477776
4	17	0.7	20	13	A2818055
5	17	0.7	31	11	N25903
6	17	0.7	37	2	HSM002040
7	17	0.7	37	11	BS175511
8	17	0.7	39	10	AM248768
9	17	0.7	44	2	HSM009691
10	17	0.7	45	2	HSM003634
11	17	0.7	49	11	BF017790
12	17	0.7	50	2	HSM003660

13	16	0.6	19	13	A2853220
14	16	0.6	19	13	A2856873
15	16	0.6	19	13	A2950028
16	16	0.6	20	13	A2370699
17	16	0.6	20	13	A2406839
18	16	0.6	21	13	A2485791
19	16	0.6	22	2	HSM002940
20	16	0.6	22	13	A2309907
21	16	0.6	22	13	TA3864070
22	16	0.6	23	13	A2390689
23	16	0.6	24	13	A2812579
24	16	0.6	25	2	HSM001398
25	16	0.6	25	2	HSM003169
26	16	0.6	25	2	HSM003450
27	16	0.6	25	13	A2832800
28	16	0.6	25	13	A2837511
29	16	0.6	25	13	TA324E07P
30	16	0.6	26	2	HSM001401
31	16	0.6	26	2	HSM001421
32	16	0.6	26	2	HSM002179
33	16	0.6	26	2	HSM002201
34	16	0.6	26	2	HSM003490
35	16	0.6	26	2	HSM003529
36	16	0.6	26	2	HSM003586
37	16	0.6	27	2	HSM001542
38	16	0.6	27	2	HSM002015
39	16	0.6	27	2	HSM002125
40	16	0.6	27	2	HSM002148
41	16	0.6	27	2	HSM003588
42	16	0.6	27	2	HSM003597
43	16	0.6	27	10	AV741507
44	16	0.6	27	11	N29432
45	16	0.6	27	11	R31539

ALIGNMENTS

RESULT	1
R71941	
LOCUS	R71941
DEFINITION	R71941 40 bp mRNA
ACCESSION	YJ84a06.r1 Soares breast 2NDHst Homo sapiens cDNA clone
VERSION	IMAGE:155410.5' similar to SP:448528 S36556; MEMBRANE GLYCOPROTEIN
KEYWORDS	CLA-1 PROTEIN LONG FORM PRECURSOR - ; mRNA sequence.
SOURCE	R71941
ORGANISM	R71941.1 GI:845973
REFERENCE	human.
AUTHORS	Homo sapiens
TITLE	Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
JOURNAL	Mammalia; Eutheria; Primates; Catarrhini; Homnidae; Homo.
COMMENT	1 (bases 1 to 40) Hillier, L., Clark, N., Dubuque, T., Elliston, K., Hawkins, M., Holman, M., Hultman, M., Kucaba, T., Le, M., Lennon, G., Mair, M., Parsons, J., Rifkin, L., Rohlfing, T., Soares, M., Tan, F., Trevisan, E., Waterston, R., Williamson, A., Wohlmann, P. and Wilson, R. The WashU-Werck EST Project Unpublished (1995) Contact: Wilson RK Washington University School of Medicine 4444 Forest Park Parkway, Box 8501, St. Louis, MO 63108 Tel: 314 286 1800 Fax: 314 286 1810 Email: est@wustl.edu Insert Size: 2714 High quality sequence starts: 1 High quality sequence stops: 1 Source: IMAGE Consortium, LNL This clone is available royalty-free through LNL: contact the IMAGE Consortium (info@image.lnl.gov) for further information. Trace considered overall poor quality Possible reversed clone: similarity on wrong strand Insert Length: 2714 Std Error: 0.00

Seq primer: M13RPI
High quality sequence stop: 1.
Location/Qualifiers

1.40
/organism="Homo sapiens"
/db_xref="GDB:573028"
/db_xref="taxon:9606"
/clone="IMAGE:155410"
/clone.lib="Scars breast 2NDBast"
/sex="Female"
/dev_stage="adult"
/lab_host="DH10B (ampicillin resistant)"
/note="Organ: breast; Vector: pT73 (Pharmacia) with a modified polylinker; Site 1: Not I; Site 2: Eco RI; 1st strand cDNA was primed with a Not I - oligo(dT) primer [5' TCTTACCATCTGCAAGTGGAGCGGCGCCCTTTTCTTTTCTTTT 3'], double-stranded cDNA was ligated to Eco RI adaptors (Pharmacia), digested with Not I and cloned into the Not I and Eco RI sites of a modified pT73 vector (Pharmacia). Library went through one round of normalization to a Cot = 230. Library constructed by Bento Soares and M.Fatima Bonaldo."

BASE COUNT 9 a 9 c 12 g 9 t 1 others
ORIGIN

Query Match 1.3%; Score 34; DB 11; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.49;
Matches 34; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 937 gtacaaggagtcagggtgttgaagcattcccc 960
|||||
Db 1 GTACAAGAGCTCAGGGGTGTTGAGCGCATCCCC 34

RESULT = 2

AZ962226

LOCUS

2M0231A02F Mouse 10kb plasmid UUGC2M library Mus musculus genomic

clone UUGC2M0231A02 F, DNA sequence.

ACCESSION AZ962226

VERSION A2962226.1

KEYWORDS GI:13833453

SOURCE GSS.

ORGANISM house mouse.

Mus musculus

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.

1 (bases 1 to 19)

Dunn,D., Aoyagi,A., Barber,M., Beacorn,T., Duval,B., Hamill,C.,

Islam,H., Longacre,S., Mahmoud,M., Meenen,E., Pedersen,T., Reilly

,M., Rose,M., Rose,R., Stokes,R., Tingey,A., von Niederhausern,A.

and Wright,D., Weiss,R.

Mouse whole genome scaffolding with paired end reads from 10kb

plasmid inserts

Unpublished (2000)

Contact: Robert B. Weiss

University of Utah Genome Center

University of Utah

Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT

84112, USA

Tel: 801 585 5606

Fax: 801 585 7177

Email: ddunne@genetics.utah.edu

Insert length: 10000 Std Error: 0.00

Plate: 0231 row: A column: 02

Seq primer: CGTTGTAAACGACGGCCACT

Class: plasmid ends

High quality sequence stop: 19.

Location/Qualifiers

1..19

/organism="Mus musculus"

/strain="C57BL/6J"

/db_xref="taxon:10090"

/clone="UUGC2M0231A02"

/clone.lib="Mouse 10kb plasmid UUGC2M library"

/sex="Female"

/lab_host="E. coli strain XL10-Gold, T1-resistant, F-"

/note="Vector: PMD42nv; Purified genomic DNA from M.

musculus C57BL/6J (Female) was obtained from the Jackson

Laboratory Mouse DNA Resource

(http://www.jax.org/resources/documents/dnares/). The DNA

was hydrodynamically sheared by repeated passage through a

0.005 inch orifice at constant velocity. The sheared DNA

was blunt end-repaired with T4 DNA polymerase and T4

polynucleotide kinase. Adaptor oligonucleotides were

ligated to the blunt ends in high molar excess. The

adapted DNA was purified and size-selected for a 9.5 to

10.5 kb range using preparative agarose gel

electrophoresis. Vector DNA was prepared from a derivative

of PMD42 (g11473211419b1AF129022.1), a copy-number

inducible derivative of plasmid R1. The vector was ligated

with adaptors complementary to the insert adaptors and

purified. The sheared, adapted mouse DNA was annealed to

adapted vector DNA, and transformed into

chemically-competent E. coli XL10-Gold (Stratagene) cells

and selected for ampicillin resistance."

BASE COUNT 16 a 0 c 2 g 1 t

ORIGIN

Query Match 0.7%; Score 19; DB 13; Length 19;
Best Local Similarity 100.0%; Pred. No. 1.1e+05;
Matches 19; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 2544 aaaaaatggaataaaaaa 2562
|||||
Db 1 AAAAAATGGAATAAAAAA 19

RESULT 3

AZ477776

LOCUS

1M0297124F Mouse 10kb plasmid UUGC1M library Mus musculus genomic

clone UUGC1M0297124 F, DNA sequence.

ACCESSION AZ477776

VERSION A2477776.1

KEYWORDS GI:10636030

SOURCE GSS.

ORGANISM house mouse.

Mus musculus

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.

1 (bases 1 to 48)

Dunn,D., Aoyagi,A., Barber,M., Beacorn,T., Duval,B., Hamill,C.,

Islam,H., Longacre,S., Mahmoud,M., Meenen,E., Pedersen,T., Reilly

,M., Rose,M., Rose,R., Stokes,R., Tingey,A., von Niederhausern,A.

and Wright,D., Weiss,R.

Mouse whole genome scaffolding with paired end reads from 10kb

plasmid inserts

Unpublished (2000)

Contact: Robert B. Weiss

University of Utah Genome Center

University of Utah

Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT

84112, USA

Tel: 801 585 5606

Fax: 801 585 7177

Email: ddunne@genetics.utah.edu

Insert length: 10000 Std Error: 0.00

Plate: 0297 row: L column: 24

Seq primer: CGTTGTAAACGACGGCCACT

Class: plasmid ends

High quality sequence stop: 48.

Location/Qualifiers

1..48

/organism="Mus musculus"

/strain="C57BL/6J"

/db_xref="taxon:10090"
 /clone="UUGC1M0297L24"
 /clone_1lb="Mouse 10kb plasmid UUGC1M library"
 /sex="Male"
 /lab_host="E. Coli strain XL10-Gold, T1-resistant, F-"
 /note="Vector: PMD42nv; Purified genomic DNA from M. musculus C57BL/6J (male) was obtained from the Jackson Laboratory Mouse DNA Resource
 (http://www.jax.org/resources/documents/dnares/). The DNA was hydrodynamically sheared by repeated passage through a 0.005 inch orifice at constant velocity. The sheared DNA was blunt end-repaired with T4 DNA polymerase and T4 polynucleotide kinase. Adaptor oligonucleotides were ligated to the blunt ends in high molar excess. The adapted DNA was purified and size-selected for a 9.5 to 10.5 kb range using preparative agarose gel electrophoresis. Vector DNA was prepared from a derivative of PMD42 (g11473211419b1AF129072.1), a copy-number inducible derivative of plasmid R1. The vector was ligated with adaptors complementary to the insert adaptors and purified. The sheared, adapted mouse DNA was annealed to adapted vector DNA, and transformed into chemically-competent E. coli XL10-Gold (Stratagene) cells and selected for ampicillin resistance."

BASE COUNT
 ORIGIN
 2 a 12 c 3 g 31 t

Query Match 0.7% Score 19; DB 13; Length 48;
 Best Local Similarity 100.0%; Pred. No. 5.8e+04;
 Matches 19; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2547 aatggaagaaagaaagaa 2565
 ||||||||||||||||||
 Db 48 AATGCAAAAAAAAAAAAAA 30

RESULT 4
 LOCUS A2818055 20 bp DNA GSS 20-FEB-2001
 DEFINITION 2M0087B23R Mouse 10kb plasmid UUGC1M library Mus musculus genomic
 A2818055
 ACCESSION A2818055
 VERSION A2818055.1 GI:12987963
 KEYWORDS GSS.
 SOURCE house mouse.
 ORGANISM Mus musculus
 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Mus;
 1 (bases 1 to 20)
 Dunn, D., Aoyagi, A., Barber, M., Beacorn, T., Duval, B., Hamil, C.,
 Islam, H., Longacre, S., Mahmoud, M., Meenen, E., Pedersen, T., Reilly,
 M., Rose, M., Rose, R., Stokes, R., Tingey, A., von Niederhausern, A.,
 and Wright, D., Weiss, R.
 Mouse whole genome scaffolding with paired end reads from 10kb
 plasmid inserts
 Unpublished (2000)
 Contact: Robert B. Weiss
 University of Utah Genome Center
 University of Utah
 Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT
 84112, USA
 Tel: 801 585 5606
 Fax: 801 585 7177
 Email: ddunn@genetics.utah.edu
 Insert length: 10000 Std Error: 0.00
 Plate: 0087 row: B column: 23
 Seq primer: CACACAGCAACACCTATGACC
 Class: plasmid ends
 High quality sequence stop: 20.
 Location/Qualifiers
 1..20
 /organism="Mus musculus"

/strain="C57BL/6J"
 /db_xref="taxon:10090"
 /clone="UUGC2M0087B23"
 /clone_1lb="Mouse 10kb plasmid UUGC1M library"
 /sex="Male"
 /lab_host="E. Coli strain XL10-Gold, T1-resistant, F-"
 /note="Vector: PMD42nv; Purified genomic DNA from M. musculus C57BL/6J (male) was obtained from the Jackson Laboratory Mouse DNA Resource
 (http://www.jax.org/resources/documents/dnares/). The DNA was hydrodynamically sheared by repeated passage through a 0.005 inch orifice at constant velocity. The sheared DNA was blunt end-repaired with T4 DNA polymerase and T4 polynucleotide kinase. Adaptor oligonucleotides were ligated to the blunt ends in high molar excess. The adapted DNA was purified and size-selected for a 9.5 to 10.5 kb range using preparative agarose gel electrophoresis. Vector DNA was prepared from a derivative of PMD42 (g11473211419b1AF129072.1), a copy-number inducible derivative of plasmid R1. The vector was ligated with adaptors complementary to the insert adaptors and purified. The sheared, adapted mouse DNA was annealed to adapted vector DNA, and transformed into chemically-competent E. coli XL10-Gold (Stratagene) cells and selected for ampicillin resistance."

BASE COUNT
 ORIGIN
 17 a 0 c 2 g 1 t

Query Match 0.7% Score 17; DB 13; Length 20;
 Best Local Similarity 100.0%; Pred. No. 5e+05;
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2550 tggagaaagaaagaaagaa 2566
 ||||||||||||||||||
 Db 1 TGGAAAAAAAAAAAAAAAAA 17

RESULT 5
 LOCUS N25903/c 31 bp mRNA EST 29-DEC-1995
 DEFINITION yw79e11.s1 Soares,placenta_8t0yweeks_2NBHptc09M Homo sapiens cDNA
 clone IMAGE:258476 3' similar to gb:X53463 GLUTATHIONE
 PEROXIDASE-GASTROINTESTINAL (HUMAN), mRNA sequence.
 ACCESSION N25903
 VERSION N25903.1 GI:1140251
 KEYWORDS EST.
 SOURCE human.
 ORGANISM Homo sapiens
 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 Mammalia; Eutheria; Primates; Catarrhini; Homnidae; Homo.
 1 (bases 1 to 31)
 Hillier, L., Clark, N., Dubuque, T., Elliston, K., Hawkins, M., Holman,
 M., Hultman, M., Kucaba, T., Le, M., Lennon, G., Marra, M., Parsons, J.,
 Rifkin, L., Rohlfing, T., Soares, M., Tan, F., Trevisakis, E., Waterston,
 R., Williamson, A., Wohlmann, P. and Wilson, R.
 The Washu-Merck EST project
 Unpublished (1995)
 Contact: Wilson RK
 Washington University School of Medicine
 444 Forest Park Parkway, Box 8501, St. Louis, MO 63108
 Tel: 314 286 1800
 Fax: 314 286 1810
 Email: est@watson.wustl.edu
 High quality sequence stops: 26
 Source: IMAGE Consortium, LLNL
 This clone is available royalty-free through LLNL; contact the
 IMAGE Consortium (info@image.llnl.gov) for further information.
 Seq primer: m13 -40 forward
 High quality sequence stop: 26.
 Location/Qualifiers
 1..31
 /organism="Homo sapiens"

/db_xref="GDB:3888086"
 /db_xref="taxon:9606"
 /clone_lib="Soares_Placenta_8to9weeks_2bHP8to9W"
 /dev_stage="two.Placentae: one from 8 weeks and another
 from 9 weeks post conception"
 /lab_host="DH10B (ampicillin resistant)"
 /note="Organ: placenta; Vector: pT73D (Pharmacia) with a
 modified polylinker. Site_1: Not I; Site_2: Eco RI; 1st
 strand cDNA was primed with a Not I - oligo(dT) primer [5'
 TGTACCAATCTGAGGAGCGCGCGCATTTTCTTTTCTTTT 3']
 double-stranded cDNA was size selected, ligated to Eco RI
 adapters (Pharmacia), digested with Not I and cloned into
 the Not I and Eco RI sites of a modified pT73 vector
 (Pharmacia). Library constructed by Bento Soares and
 M.Falima Bernaldo."

BASE COUNT 6 a 5 c 0 g 20 t

ORIGIN

Query Match 0.7%; Score 17; DB 11; Length 31;
 Best Local Similarity 100.0%; Pred. No. 3.7e+05;
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 2550 tggaaaaaataaaaaa 2566
 ||||||||||||||||
 Db 22 TCGAATAAAAAAAAAA 6

RESULT 6

HSMD02040/c standard; RNA; EST; 37 BP.

AC AL037709;
 XX
 SV AL037709.1

DT 12-MAR-1999 (Rel. 59, Created)
 DT 12-MAR-1999 (Rel. 59, Last updated, Version 1)

DE Homo sapiens mRNA; EST DKFZP564A157_s1 (from clone DKFZP564A157)

XX EST; expressed sequence tag.

OS Homo sapiens (human)

OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
 CC Eutheria; Primates; Catarrhini; Homidae; Homo.

XX [1]

RP 1-37

RA Bloeker H., Boeher M., Brandt P., Mewes W., Gassenhuber J., Wiemann S.;

RT Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.

RL MIPS, Am Kopfersplitz 18a D-82152 Martinsried, GERMANY

CC Clone from S. Wiemann, sequenced by GBF within the cDNA

CC sequencing consortium of the German Genome Project

CC No. 1 sequence available

CC This clone is available at the RZPD in Berlin

CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059

CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de

XX key Location/Qualifiers

FT source 1. 37
 FT /db_xref="taxon:9606"
 FT /organism="Homo sapiens"
 FT /clone="DKFZP564A157"
 FT /clone_lib="564 (synonym: hfbz2). Vector pAMP1; host
 FT X1-2blue; sites NotI + SalI"
 FT /dev_stage="fetal"
 FT /tissue_type="brain"
 XX

SO Sequence 37 BP; 4 A; 10 C; 0 G; 23 T; 0 other;

Query Match 0.7%; Score 17; DB 2; Length 37;
 Best Local Similarity 100.0%; Pred. No. 3.3e+05;
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 2550 tggaaaaaataaaaaa 2566
 ||||||||||||||||
 Db 22 TCGAATAAAAAAAAAA 6

RESULT 7

BG175511 37 bp mRNA EST 06-FEB-2001

LOCUS 602334778F2 NCL_CGAP_Mam1 Mus musculus cDNA clone IMAGE:4457995 5',

DEFINITION mRNA sequence.

ACCESSION BG175511 GI:12682214

VERSION BG175511.1

KEYWORDS EST.

SOURCE house mouse.

ORGANISM Mus musculus

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.

REFERENCE 1 (bases 1 to 37)

NIH-MGC http://mgs.nci.nih.gov/
 National Institutes of Health, Mammalian Gene Collection (MGC)

Unpublished (1999)

Contact: Robert Strausberg, Ph.D.
 Email: cga@rs-femail.nih.gov

Tissue Procurement: Gilbert Smith, Ph.D.
 cDNA Library Preparation: Life Technologies, Inc.

cDNA Library Arrayed by: The I.M.A.G.E. Consortium (LNL)

DNA sequencing by: Incyte Genomics, Inc.

Clone distribution: MGC clone distribution information can be
 found through the I.M.A.G.E. Consortium/LNL at:

http://image.lnl.gov

Plate: LLM10255 row: k column: 20

High quality sequence stop: 32.

FEATURES

source 1. 37

/organism="Mus musculus"

/strain="FVB/N"

/db_xref="taxon:10090"

/clone="IMAGE:4457995"

/clone_lib="NCL_CGAP_Mam1"

/tissue_type="tumor, biopsy sample"

/dev_stage="3 months, virgin"

/lab_host="DH10B"

/note="Organ: mammary; Vector: pCMV-SPORT6; Site_1: SalI;
 Site_2: NotI; Cloned unidirectionally. Primer: Oligo dT.
 Library constructed by Life Technologies. Investigator
 providing samples: Gilbert Smith, NIH"

BASE COUNT 23 a 2 c 5 g 7 t

ORIGIN

Query Match 0.7%; Score 17; DB 11; Length 37;
 Best Local Similarity 100.0%; Pred. No. 3.3e+05;
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 2550 tggaaaaaataaaaaa 2566
 ||||||||||||||||
 Db 9 TCGAATAAAAAAAAAA 25

RESULT 8

AM248768 39 bp mRNA EST 07-JAN-2000

LOCUS 2820919.3prime NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2820919 3',

DEFINITION mRNA sequence.

ACCESSION AM248768

VERSION AM248768.1 GI:6591761


```

KEYWORDS EST.
SOURCE human.
ORGANISM Homo sapiens
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Homiidae; Homo.
REFERENCE 1 (bases 1 to 39)
AUTHORS NIH-MGC http://mgi.nci.nih.gov/.
TITLE National Institutes of Health, Mammalian Gene Collection (MGC)
JOURNAL Unpublished (1999)
COMMENT Other_ESTS: 2820919.Sprime.
Contact: Robert Strausberg, Ph.D.
Email: cgabbs-r@mail.nih.gov
Tissue Procurement: DCTD/DTF CDNA Library Preparation: Ling
Hong/Rubin Laboratory CDNA Library Arrayed by: The I.M.A.G.E.
Consortium (LNL) DNA Sequencing by: Berkeley MGC sequencing
project Clone distribution: MGC clone distribution information can
be found through the I.M.A.G.E. Consortium/LNL at:
www.bio.lnl.gov/bdrr/image/image.html Base Calling / Quality
Scores: PHRED from University of Washington Genome Center. Vector
Trimming: cross-match from University of Washington Genome Center
PHRAP suite. Poly-T identification: patmatch.pl from Berkeley
Drosophila Genome Project. University of Washington Genome Center:
http://www.genome.washington.edu Low Quality Sequence: 7 contiguous
PHRED high quality bases following vector sequence. Very low
Quality Sequence: Trace file contained 39 contiguous distinct peaks
following vector sequence. Polyadenylation: Based upon the presence
of a xhoi site followed by a run of 14 or more T residues at the
beginning of the sequence, this cDNA insert was polyadenylated.
Plate: LNCM5 row: H column: 8
High quality sequence stop: 7.
Location/Qualifiers
1. 39
source
/organism="Homo sapiens"
/db_xref="taxon:9606"
/clone_lib="IMAGE:2820919"
/clone_lib="NIH-MGC-7"
/tissue_type="small cell carcinoma"
/cell_line="MGC3"
/lab_host="DH10B (phage-resistant)"
/notes="Organ: lung; Vector: pOTB7; Site_1: XhoI; Site_2:
EcoRI; cDNA made by oligo-dt priming. Directionally
cloned into EcoRI/XhoI sites using the following 5'
adaptor: GGCACGAG(G). Size-selected >500bp for average
insert size 1.8kb. Library constructed by Ling Hong in
the laboratory of Gerald M. Rubin (University of
California, Berkeley) using ZAP-cDNA synthesis kit
(Stratagene) and Superscript II RT (Life Technologies)."
```

```

BASE COUNT 6 a 2 c 1 g 30 t
ORIGIN
Query Match 0.7%; Score 17; DB 10; Length 39;
Best Local Similarity 100.0%; Pred. No. 3.2e+05;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 2550 tggagaaaaaaaaaaaaa 2566
|||||
Db 34 TGGAAAAA 18

RESULT 9
HSM009691/c
ID HSM009691 standard; RNA; EST; 44 BP.
AC AL044841;
XX
SV AL044841.1
XX
DT 12-MAR-1999 (Rel. 59, Created)
DT 12-MAR-1999 (Rel. 59, Last updated, Version 1)
XX
DE Homo sapiens mRNA; EST DKFZp434B083_s1 (from clone DKFZp434B083)
XX

KM EST; expressed sequence tag.
XX
XX Homo sapiens (human)
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
OC Eutheria; Primates; Catarrhini; Homiidae; Homo.
XX
XX [1]
RP 1-44
RA Mambuth R., Heubner D., Mewes W., Gassenhuber J., Wiemann S.;
RT Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
RL MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
XX
CC Clone from S. Wiemann, sequenced by AGOMA within the CDNA
CC sequencing consortium of the German Genome Project
CC This clone is available at the RZPD in Berlin
CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
XX
FH Key Location/Qualifiers
FH source 1. 44
FT /db_xref="taxon:9606"
FT /organism="Homo sapiens"
FT /clone_lib="DKFZp434B083"
FT /clone_lib="434 (synonym: htes3). Vector psport1; host
FT DH10B; sites NotI + SalI"
FT /dev_stage="adult"
FT /tissue_type="testis"
XX
Sequence 44 BP; 6 A; 8 C; 3 G; 27 T; 0 other;
Query Match 0.7%; Score 17; DB 2; Length 44;
Best Local Similarity 100.0%; Pred. No. 3e+05;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 2550 tggagaaaaaaaaaaaaa 2566
|||||
Db 38 TGGAAAAA 22

RESULT 10
HSM003634/c
ID HSM003634 standard; RNA; EST; 45 BP.
AC AL039158;
XX
SV AL039158.1
XX
DT 12-MAR-1999 (Rel. 59, Created)
DT 12-MAR-1999 (Rel. 59, Last updated, Version 1)
XX
DE Homo sapiens mRNA; EST DKFZp566M244_s1 (from clone DKFZp566M244)
XX
XX EST; expressed sequence tag.
XX
XX Homo sapiens (human)
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
OC Eutheria; Primates; Catarrhini; Homiidae; Homo.
XX
XX [1]
RP 1-45
RA Bloecher H., Boecher M., Brandt P., Mewes W., Gassenhuber J., Wiemann S.;
RT Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
RL MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
XX
CC Clone from S. Wiemann, sequenced by GBF within the CDNA
CC sequencing consortium of the German Genome Project
CC No r1 sequence available
CC This clone is available at the RZPD in Berlin
```

CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
 CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
 XX
 FH Key Location/Qualifiers
 FT source 1. 45
 FT /db_xref="taxon:9606"
 FT /organism="Homo sapiens"
 FT /clone="DKFZ566M244"
 FT /clone_lib="566 (synonym: hfk2). Vector pAMP1; host
 FT X1-2blue; sites NotI + SalI"
 FT /dev_stage="fetal"
 FT /tissue_type="kidney"
 FT
 XX
 SQ Sequence 45 BP; 2 A; 10 C; 0 G; 33 T; 0 other;
 Query Match 0.7%; Score 17; DB 2; Length 45;
 Best Local Similarity 100.0%; Pred. No. 2.9e+05;
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 2550 tggaaaaaaaaaaaaa 2566
 Db 37 TGGAAAAAAAAAAAAA 21
 RESULT 11
 LOCUS BF017790 49 bp mRNA EST 29-DEC-2000
 DEFINITION ux75h05.y1 McCarrey Eddy type B spermatogonia Mus musculus cDNA
 ACCESSION BF017790
 VERSION BF017790
 KEYWORDS EST.
 SOURCE house mouse.
 ORGANISM Mus musculus
 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
 1 (bases 1 to 49)
 AUTHORS Marra, M., Hillier, L., Kucaba, T., Martin, J., Beck, C., Wylie, T.,
 Underwood, K., Stepien, M., Theising, B., Allen, M., Bowers, Y., Person
 E., Kohn, S., Shih, T., Jackson, Y., Cardenas, M., McCann, R.,
 Waterston, R., and Wilson, R.
 The WashU-NCI Mouse EST Project 1999
 Unpublished (1999)
 CONTACT: Marra M/WashU-NCI Mouse EST Project 1999
 Washington University School of Medicine
 4444 Forest Park Parkway, Box 8501, St. Louis, MO 63108, USA
 Tel: 314 286 1800
 Fax: 314 286 1810
 Email: mouseest@wustl.edu
 This clone is available royalty-free through LNL; contact the
 IMAGE Consortium (info@image.lnl.gov) for further information.
 WGI:141697
 Seq primer: Primer name ambiguous.
 FEATURES
 source Location/Qualifiers
 1. 49
 /organism="Mus musculus"
 /strain="CD-1"
 /db_xref="taxon:10090"
 /clone="IMAGE:3654393"
 /clone_lib="McCarrey Eddy type B spermatogonia"
 /sex="male"
 /tissue_type="type B spermatogonia, pooled from multiple
 mice"
 /dev_stage="8 day"
 /lab_host="DH10B (phage-resistant)"
 /note="Organ: testis; Vector: pBluescript SK+ (Stratagene
); Site_1: XhoI; Site_2: EcoRI; cDNA oligo df-primed
 [5'-(GA)10-ACTAGCTCGAGTTTGTTTT-3'] and directionally
 cloned using 5' linkers 5'-AATTGGCGACGAG-3' and
 5'-CTGCTGCCG-3'. Size selection of >400bp material gives

average insert size ranging from 1-2 kb. Library was mass
 excised (from lambda-UniZAP-XR) and resulting
 single-stranded phagemids were prepped and transformed
 into DH10B. Library contains 968 recombinants.
 References: J. Androl. 20:635-639 and Gene 23:263-269.
 Library constructed and donated by J. McCarrey, Ph.D.
 (Southwest Foundation for Biomedical Research, Dept. of
 Genetics); excision done by E.M. Eddy, Ph.D. (National
 Institutes of Health, National Institute of Environmental
 Health Sciences). Original lambda-instant library is
 available through ATCC, catalog #63417.
 BASE COUNT 24 a 9 c 9 g 7 t
 ORIGIN
 Query Match 0.7%; Score 17; DB 11; Length 49;
 Best Local Similarity 100.0%; Pred. No. 2.8e+05;
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 2550 tggaaaaaaaaaaaaa 2566
 Db 24 TGGAAAAAAAAAAAAA 40
 RESULT 12
 HSM003660/c standard; RNA; EST; 50 BP.
 ID HSM003660
 AC AL039184;
 SV AL039184.1
 DT 12-MAR-1999 (Rel. 59, Created)
 DT 12-MAR-1999 (Rel. 59, Last updated, Version 1)
 XX
 DE Homo sapiens mRNA; EST DKFZ5660064.s1 (from clone DKFZ5660064)
 XX
 KW EST; expressed sequence tag.
 OS Homo sapiens (human)
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
 OC Eutheria; Primates; Catarrhini; Hominiidae; Homo.
 XX
 RN [1]
 RP 1-50
 RA Bloecher H., Boecher M., Brandt P., Mewes W., Gassenhuber J., Wiemann S.;
 Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
 RL MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
 CC
 CC Clone from S. Wiemann, sequenced by GBR within the cDNA
 CC sequencing consortium of the German Genome Project
 CC No r1 sequence available
 CC This clone is available at the RZPD in Berlin
 CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
 CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
 XX
 FH Key Location/Qualifiers
 1. 50
 /db_xref="taxon:9606"
 /organism="Homo sapiens"
 /clone="DKFZ5660064"
 /clone_lib="566 (synonym: hfk2). Vector pAMP1; host
 FT X1-2blue; sites NotI + SalI"
 FT /dev_stage="fetal"
 FT /tissue_type="kidney"
 FT
 XX
 SQ Sequence 50 BP; 10 A; 10 C; 2 G; 28 T; 0 other;
 Query Match 0.7%; Score 17; DB 2; Length 50;
 Best Local Similarity 100.0%; Pred. No. 2.7e+05;

Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 2550 tggaaaaaataaaaaa 2566
 ||||||||||||||||
 Db 21 TCGAAAAAATAAAAAA 5

RESULT 13

AZ853220 19 bp DNA GSS 21-FEB-2001
 LOCUS 2M016J15F Mouse 10kb plasmid UUGC1M library Mus musculus genomic
 DEFINITION clone UUGC2M016J15 F, DNA sequence.
 ACCESSION AZ853220
 VERSION AZ853220.1 GI:13041116
 KEYWORDS GSS.
 SOURCE house mouse.
 ORGANISM Mus musculus

REFERENCE

AUTHORS Dunn,D., Aoyagi,A., Barber,M., Beacorn,T., Duval,B., Hamil,C.,
 Islam,H., Longacre,S., Mahmoud,M., Meenen,E., Pedersen,T., Rellily,
 M., Rose,M., Rose,R., Stokes,R., Tingey,A., von Niederhausern,A.
 and Wright,D., Weiss,R.
 Mouse whole genome scaffolding with paired end reads from 10kb
 plasmid inserts
 Unpublished (2000)

JOURNAL

COMMENT Contact: Robert B. Weiss
 University of Utah Genome Center
 University of Utah
 Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT
 84112, USA
 Tel: 801 585 5606
 Fax: 801 585 7177
 Email: dunn@genetics.utah.edu
 Insert Length: 10000 Std Error: 0.00
 Plate: 0156 row: J column: 15
 Seq primer: CGTTGTAACGACGCGCCACT
 Class: plasmid ends
 High quality sequence stop: 19.

FEATURES

Location/Qualifiers

1..19
 /organism="Mus musculus"
 /strain="C57BL/6J"
 /db_xref="taxon:10090"
 /clone="UUGC2M016J15"
 /clone_1lb="Mouse 10kb plasmid UUGC1M library"
 /sex="Male"
 /lab_host="E. Coli strain XL10-Gold, TI-resistant, F-"
 /note="Vector: PMD42nv. Purified genomic DNA from M.
 musculus C57BL/6J (male) was obtained from the Jackson
 Laboratory Mouse DNA Resource
 (http://www.jax.org/resources/documents/dnares/). The DNA
 was hydrodynamically sheared by repeated passage through a
 0.005 inch orifice at constant velocity. The sheared DNA
 was blunt end-repaired with T4 DNA polymerase and T4
 polynucleotide kinase. Adaptor oligonucleotides were
 ligated to the blunt ends in high molar excess. The
 adapted DNA was purified and size-selected for a 9.5 to
 10.5 kb range using preparative agarose gel
 electrophoresis. Vector DNA was prepared from a derivative
 of PMD42 (g114732114[gb|AF129072.1]), a copy-number
 inducible derivative of plasmid R1. The vector was ligated
 with adaptors complementary to the insert adaptors and
 purified. The sheared, adapted mouse DNA was annealed to
 adapted vector DNA, and transformed into
 chemically-competent E. coli XL10-Gold (Stratagene) cells
 and selected for ampicillin resistance."

BASE COUNT

17 a 0 c 2 g 0 t

Query Match

0.6%; Score 16; DB 13; Length 19;

Best Local Similarity 100.0%; Pred. No. 1.1e+06;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 2551 ggaataaaaaataaaaa 2566
 ||||||||||||||||
 Db 1 GGAATAAATAAATAAATA 16

RESULT 14

AZ856873 19 bp DNA GSS 21-FEB-2001
 LOCUS 2M016J19F Mouse 10kb plasmid UUGC1M library Mus musculus genomic
 DEFINITION clone UUGC2M016J19 F, DNA sequence.
 ACCESSION AZ856873
 VERSION AZ856873.1 GI:13048296
 KEYWORDS GSS.
 SOURCE house mouse.
 ORGANISM Mus musculus

REFERENCE

AUTHORS Dunn,D., Aoyagi,A., Barber,M., Beacorn,T., Duval,B., Hamil,C.,
 Islam,H., Longacre,S., Mahmoud,M., Meenen,E., Pedersen,T., Rellily,
 M., Rose,M., Rose,R., Stokes,R., Tingey,A., von Niederhausern,A.
 and Wright,D., Weiss,R.
 Mouse whole genome scaffolding with paired end reads from 10kb
 plasmid inserts
 Unpublished (2000)

JOURNAL

COMMENT Contact: Robert B. Weiss
 University of Utah Genome Center
 University of Utah
 Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT
 84112, USA
 Tel: 801 585 5606
 Fax: 801 585 7177
 Email: dunn@genetics.utah.edu
 Insert Length: 10000 Std Error: 0.00
 Plate: 0161 row: O column: 19
 Seq primer: CGTTGTAACGACGCGCCACT
 Class: plasmid ends
 High quality sequence stop: 19.

FEATURES

Location/Qualifiers

1..19
 /organism="Mus musculus"
 /strain="C57BL/6J"
 /db_xref="taxon:10090"
 /clone="UUGC2M016J19"
 /clone_1lb="Mouse 10kb plasmid UUGC1M library"
 /sex="Male"
 /lab_host="E. Coli strain XL10-Gold, TI-resistant, F-"
 /note="Vector: PMD42nv. Purified genomic DNA from M.
 musculus C57BL/6J (male) was obtained from the Jackson
 Laboratory Mouse DNA Resource
 (http://www.jax.org/resources/documents/dnares/). The DNA
 was hydrodynamically sheared by repeated passage through a
 0.005 inch orifice at constant velocity. The sheared DNA
 was blunt end-repaired with T4 DNA polymerase and T4
 polynucleotide kinase. Adaptor oligonucleotides were
 ligated to the blunt ends in high molar excess. The
 adapted DNA was purified and size-selected for a 9.5 to
 10.5 kb range using preparative agarose gel
 electrophoresis. Vector DNA was prepared from a derivative
 of PMD42 (g114732114[gb|AF129072.1]), a copy-number
 inducible derivative of plasmid R1. The vector was ligated
 with adaptors complementary to the insert adaptors and
 purified. The sheared, adapted mouse DNA was annealed to
 adapted vector DNA, and transformed into
 chemically-competent E. coli XL10-Gold (Stratagene) cells
 and selected for ampicillin resistance."

BASE COUNT

0 a 5 c 0 g 14 t

ORIGIN

Query Match 0.6%; Score 16; DB 13; Length 19;
Best Local Similarity 100.0%; Pred. No. 1.1e+06;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaaaaa 2566
Db 16 GGAATAAAAAAAAA 1

RESULT 15

AZ950028

LOCUS 19 bp DNA GSS 27-APR-2001
DEFINITION 2M0213L19R Mouse 10kb plasmid UUGC2M library Mus musculus genomic
clone UUGC2M0213L19 R. DNA sequence.

ACCESSION

AZ950028

VERSION AZ950028.1 GI:13821255

KEYWORDS

GSS.

SOURCE

house mouse.

ORGANISM

Unpublished (2000)
Contact: Robert B. Weiss
University of Utah Genome Center
University of Utah
Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLIC, UT
84112, USA
Tel: 801 585 5606
Fax: 801 585 7177
Email: ddunn@genetics.utah.edu
Insert Length: 10000 Std Error: 0.00
Plate: 0213 row: L column: 19
Seq primer: CACACAGAAACAGCTATGACC
Class: plasmid ends
High quality sequence stop: 19.

TITLE

Unpublished (2000)

JOURNAL

COMMENT

COMMENT

FEATURES

source

1. 19

Location/Qualifiers

/organism="Mus musculus"

/strain="C57BL/6J"

/db_xref="taxon:10090"

/clone="UUGC2M0213L19"

/clone_lib="Mouse 10kb plasmid UUGC2M library"

/sex="Female"

/lab_host="E. coli strain XL10-Gold, TI-resistant, F-"

/note="Vector: PMD42nv. Purified genomic DNA from M. musculus C57BL/6J (female) was obtained from the Jackson Laboratory Mouse DNA Resource (http://www.jax.org/resources/documents/dnares/). The DNA was hydrodynamically sheared by repeated passage through a 0.005 inch orifice at constant velocity. The sheared DNA was blunt end-repaired with T4 DNA polymerase and T4 polynucleotide kinase. Adaptor oligonucleotides were ligated to the blunt ends in high molar excess. The adapted DNA was purified and size-selected for a 9.5 to 10.5 kb range using preparative agarose gel electrophoresis. Vector DNA was prepared from a derivative of PMD42 (g114732114[9b]AF129072.1), a copy-number inducible derivative of plasmid R1. The vector was ligated with adaptors complementary to the insert adaptors and purified. The sheared, adapted mouse DNA was annealed to adapted vector DNA, and transformed into chemically-competent E. coli XL10-Gold (Stratagene) cells and selected for ampicillin resistance."

Query Match 0.6%; Score 16; DB 13; Length 19;
Best Local Similarity 100.0%; Pred. No. 1.1e+06;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaaaaa 2566
Db 4 GGAATAAAAAAAAA 19

RESULT 16

AZ370699/c

LOCUS 20 bp DNA GSS 02-OCT-2000
DEFINITION 1M0121N17R Mouse 10kb plasmid UUGC1M library Mus musculus genomic
clone UUGC1M0121N17 R. DNA sequence.

ACCESSION

AZ370699

VERSION AZ370699.1 GI:10484399

KEYWORDS

GSS.

SOURCE

house mouse.

ORGANISM

Unpublished (2000)
Contact: Robert B. Weiss
University of Utah Genome Center
University of Utah
Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLIC, UT
84112, USA
Tel: 801 585 5606
Fax: 801 585 7177
Email: ddunn@genetics.utah.edu
Insert Length: 10000 Std Error: 0.00
Plate: 0121 row: N column: 17
Seq primer: CACACAGAAACAGCTATGACC
Class: plasmid ends
High quality sequence stop: 20.

TITLE

Unpublished (2000)

JOURNAL

COMMENT

COMMENT

FEATURES

source

1. 20

Location/Qualifiers

/organism="Mus musculus"

/strain="C57BL/6J"

/db_xref="taxon:10090"

/clone="UUGC1M0121N17"

/clone_lib="Mouse 10kb plasmid UUGC1M library"

/sex="Male"

/lab_host="E. coli strain XL10-Gold, TI-resistant, F-"

/note="Vector: PMD42nv. Purified genomic DNA from M. musculus C57BL/6J (male) was obtained from the Jackson Laboratory Mouse DNA Resource (http://www.jax.org/resources/documents/dnares/). The DNA was hydrodynamically sheared by repeated passage through a 0.005 inch orifice at constant velocity. The sheared DNA was blunt end-repaired with T4 DNA polymerase and T4 polynucleotide kinase. Adaptor oligonucleotides were ligated to the blunt ends in high molar excess. The adapted DNA was purified and size-selected for a 9.5 to 10.5 kb range using preparative agarose gel electrophoresis. Vector DNA was prepared from a derivative of PMD42 (g114732114[9b]AF129072.1), a copy-number inducible derivative of plasmid R1. The vector was ligated with adaptors complementary to the insert adaptors and purified. The sheared, adapted mouse DNA was annealed to adapted vector DNA, and transformed into chemically-competent E. coli XL10-Gold (Stratagene) cells and selected for ampicillin resistance."

BASE COUNT

ORIGIN

14 a 0 c 5 g 0 t

BASE COUNT

ORIGIN

0 a 2 c 0 g 18 t

Query Match 0.6%; Score 16; DB 13; Length 20;

Best Local Similarity 100.0%; Pred. No. 1.1e+06;

Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 2551 ggaagaaagaaagaaagaa 2566
|||||

Db 20 GGAAAAAAGAAAAA 5

RESULT 17

AZ406839

LOCUS 20 bp DNA GSS 03-OCT-2000
DEFINITION 1M0176C16F Mouse 10kb plasmid UUGC1M library Mus musculus genomic

clone UUGC1M0176C16 F, DNA sequence.

ACCESSION AZ406839
VERSION AZ406839.1 GI:10530852

KEYWORDS GSS.

SOURCE house mouse;
ORGANISM Mus musculus

REFERENCE Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.

AUTHORS

Dunn,D., Aoyagi,A., Barber,M., Beacorn,T., Duval,B., Hamil,C.,
Islam,H., Longacre,S., Mahmoud,M., Meenen,E., Pedersen,T., Reilly,
M., Rose,M., Rose,R., Stokes,R., Tinney,A., von Niederhausern,A.
and Wright,D., Weiss,R.

TITLE Mouse whole genome scaffolding with paired end reads from 10kb
plasmid inserts

JOURNAL Unpublished (2000)

COMMENT Contact: Robert B. Weiss
University of Utah Genome Center
Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT
84112, USA
Tel: 801 585 5606
Fax: 801 585 7177
Email: ddunne@genetics.utah.edu
Insert Length: 10000 Std Error: 0.00
Plate: 0176 row: C column: 16
Seq primer: CGTTGTAAACGACGCCAGT
Class: plasmid ends
High quality sequence stop: 20.
Location/Qualifiers

1. 20
/organism="Mus musculus"
/strain="C57BL/6J"
/db_xref="taxon:10090"
/clone="UUGC1M0176C16"
/clone_lib="Mouse 10kb plasmid UUGC1M library"
/sex="Male"
/lab_host="E. Coli strain XL10-Gold, T1-resistant, F-"
/note="Vector: PWD42nv; Purified genomic DNA from M.
musculus C57BL/6J (male) was obtained from the Jackson
Laboratory Mouse DNA Resource
(http://www.jax.org/resources/documents/dnares/). The DNA
was hydrodynamically sheared by repeated passage through a
0.005 inch orifice at constant velocity. The sheared DNA
was blunt end-repaired with T4 DNA polymerase and T4
polynucleotide kinase. Adaptor oligonucleotides were
ligated to the blunt ends in high molar excess. The
adapted DNA was purified and size-selected for a 9.5 to
10.5 kb range using preparative agarose gel
electrophoresis. Vector DNA was prepared from a derivative
of PWD42 (g114732114[9b]AF129072.1), a copy-number
inducible derivative of plasmid R1. The vector was ligated
with adaptors complementary to the insert adaptors and
purified. The sheared, adapted mouse DNA was annealed to
adapted vector DNA, and transformed into
chemically-competent E. coli XL10-Gold (Stratagene) cells
and selected for ampicillin resistance."

BASE COUNT

8 a 7 g 2 g 3 t

ORIGIN

Query Match 0.6%; Score 16; DB 13; Length 20;

Best Local Similarity 100.0%; Pred. No. 1.1e+06;

Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1768 caccatcacacatg 1783
|||||

Db 3 CACACATCACACATG 18

RESULT 18

AZ785791 21 bp DNA GSS 16-FEB-2001
LOCUS AZ785791/C

DEFINITION 1M0030019F Mouse 10kb plasmid UUGC1M library Mus musculus genomic

clone UUGC1M0030019 F, DNA sequence.

ACCESSION AZ785791
VERSION AZ785791.1 GI:12922904

KEYWORDS GSS.

SOURCE house mouse;
ORGANISM Mus musculus

REFERENCE Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.

AUTHORS

Dunn,D., Aoyagi,A., Barber,M., Beacorn,T., Duval,B., Hamil,C.,
Islam,H., Longacre,S., Mahmoud,M., Meenen,E., Pedersen,T., Reilly,
M., Rose,M., Rose,R., Stokes,R., Tinney,A., von Niederhausern,A.
and Wright,D., Weiss,R.

TITLE Mouse whole genome scaffolding with paired end reads from 10kb
plasmid inserts

JOURNAL Unpublished (2000)

COMMENT Contact: Robert B. Weiss
University of Utah Genome Center
Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT
84112, USA
Tel: 801 585 5606
Fax: 801 585 7177
Email: ddunne@genetics.utah.edu
Insert Length: 10000 Std Error: 0.00
Plate: 0030 row: O column: 19
Seq primer: CGTTGTAAACGACGCCAGT
Class: plasmid ends
High quality sequence stop: 21.
Location/Qualifiers

1. 21
/organism="Mus musculus"
/strain="C57BL/6J"
/db_xref="taxon:10090"
/clone="UUGC1M0030019"
/clone_lib="Mouse 10kb plasmid UUGC1M library"
/sex="Male"
/lab_host="E. Coli strain XL10-Gold, T1-resistant, F-"
/note="Vector: PWD42nv; Purified genomic DNA from M.
musculus C57BL/6J (male) was obtained from the Jackson
Laboratory Mouse DNA Resource
(http://www.jax.org/resources/documents/dnares/). The DNA
was hydrodynamically sheared by repeated passage through a
0.005 inch orifice at constant velocity. The sheared DNA
was blunt end-repaired with T4 DNA polymerase and T4
polynucleotide kinase. Adaptor oligonucleotides were
ligated to the blunt ends in high molar excess. The
adapted DNA was purified and size-selected for a 9.5 to
10.5 kb range using preparative agarose gel
electrophoresis. Vector DNA was prepared from a derivative
of PWD42 (g114732114[9b]AF129072.1), a copy-number
inducible derivative of plasmid R1. The vector was ligated
with adaptors complementary to the insert adaptors and
purified. The sheared, adapted mouse DNA was annealed to
adapted vector DNA, and transformed into
chemically-competent E. coli XL10-Gold (Stratagene) cells
and selected for ampicillin resistance."

BASE COUNT

8 a 7 g 2 g 3 t

BASE COUNT 0 a 7 c 0 g 14 t
ORIGIN

Query Match 0.6%; Score 16; DB 13; Length 21;
Best Local Similarity 100.0%; Pred. No. 1.1e+06;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 2551 ggaataaaataaa 2566
|||||
Db 16 GGAATAAAATAAA 1

RESULT 19
HSM002940
ID HSM002940 standard; RNA; EST; 22 BP.

SV AL038464.1
XX 12-MAR-1999 (Rel. 59, Created)
DT 12-MAR-1999 (Rel. 59, Last updated, Version 1)

XX Homo sapiens mRNA; EST DKFP566B0646_r1 (from clone DKFP566B0646)

XX EST; expressed sequence tag.

XX Homo sapiens (human)
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
OC Eutheria; Primates; Catarrhini; Homidae; Homo.

XX [1]
RP 1-22
RA O'Brienaeider B., Obermaier B., Neues W., Gassenhuber J., Wiemann S.;
RT Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
RL MIPS, Am Klopferplatz 18a D-81552 Martinsried, GERMANY

XX Clone from S. Wiemann, sequenced by MedGenomix within the CDNA
CC sequencing consortium of the German Genome Project

CC s1 sequence also available
CC This clone is available at the RZPD in Berlin
CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de

XX Key Location/Qualifiers

FT source 1. 22
FT /db_xref="taxon:9606"
FT /organism="Homo sapiens"
FT /clone_id="DKFP566B0646"
FT /clone_lib="566 (synonym: hfk2). Vector pAMP1, host
FT X1-Zblue; sites: NotI + SalI"
FT /dev_stage="fetal"
FT /tissue_type="kidney"

XX Sequence 22 BP; 20 A; 0 C; 2 G; 0 T; 0 other;

Query Match 0.6%; Score 16; DB 2; Length 22;
Best Local Similarity 100.0%; Pred. No. 1e+06;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 2551 ggaataaaataaa 2566
|||||
Db 2 GGAATAAAATAAA 17

RESULT 20
AZ309907/c 22 bp DNA GSS 29-SEP-2000
LOCUS IN0017N14F Mouse 10kb plasmid UUGC1M library Mus musculus genomic

ACCESSION clone UUGC1M0017N14 F, DNA sequence.
AZ309907
VERSION 1. GI:10351367
KEYWORDS GSS.

SOURCE house mouse.
ORGANISM Mus musculus

REFERENCE
AUTHORS Dunn, D., Aoyagi, A., Barber, M., Beacorn, T., Duval, B., Hamill, C.,
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
1 (bases 1 to 22)

TITLE Mouse whole genome scaffolding, with paired end reads from 10kb
plasmid inserts

JOURNAL Contact: Robert B. Weiss
COMMENT University of Utah Genome Center
Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLIC, UT
84112, USA

Tel: 801 585 5606
Fax: 801 585 7177
Email: ddunn@genetics.utah.edu

Insert Length: 10000 Std Error: 0.00
Plate: 0017 row: N column: 14
Seq primer: CGTTGTAACGACGCCACT

Class: plasmid ends
High quality sequence stop: 22.

FEATURES
source

1. 22
Location/Qualifiers
/organism="Mus musculus"
/strain="C57BL/6J"
/db_xref="taxon:10090"
/clone="UUGC1M0017N14"
/clone_lib="Mouse 10kb plasmid UUGC1M library"
/sex="Male"
/lab_host="E. coli strain XL10-Gold, T1-resistant, F-"
/note="Vector: PMD42nv; Purified genomic DNA from M.
musculus C57BL/6J (male) was obtained from the Jackson
Laboratory Mouse DNA Resource
(http://www.jax.org/resources/documents/dnares/). The DNA
was hydrodynamically sheared by repeated passage through a
0.005 inch orifice at constant velocity. The sheared DNA
was blunt end-repaired with T4 DNA polymerase and T4
polynucleotide kinase. Adaptor oligonucleotides were
ligated to the blunt ends in high molar excess. The
adapted DNA was purified and size-selected for a 9.5 to
10.5 kb range using preparative agarose gel
electrophoresis. Vector DNA was prepared from a derivative
of PMD42 (g114732114[gb]AF129072.1), a copy-number
inducible derivative of plasmid R1. The vector was ligated
with adaptors complementary to the insert adaptors and
purified. The sheared, adapted mouse DNA was annealed to
adapted vector DNA, and transformed into
chemically-competent E. coli XL10-Gold (Stratagene) cells
and selected for ampicillin resistance."

BASE COUNT 0 a 3 c 0 g 19 t
ORIGIN

Query Match 0.6%; Score 16; DB 13; Length 22;
Best Local Similarity 100.0%; Pred. No. 1e+06;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 2551 ggaataaaataaa 2566
|||||
Db 21 GGAATAAAATAAA 6

RESULT 21
TA386H07Q/c 22 bp DNA GSS 13-DEC-2000
LOCUS TA386H07Q

DEFINITION T. brucei sheared genomic DNA clone 386h07, reverse sequence, genomic survey sequence.

ACCESSION AL498291

VERSION AL498291.1 GI:11874013

KEYWORDS GSS.

SOURCE Trypanosoma brucei.

ORGANISM Trypanosoma brucei.

Eukaryota; Euglenozoa; Kinetoplastida; Trypanosomatidae;

REFERENCE 1 (bases 1 to 22)

AUTHORS Hall, N., Bowman, S., Lennard, N.J., Doggett, J., Atkin, R., Chillingworth, C., Ormond, D., Harris, B., El-Sayed, N., Hou, L., Melville, S.E., Rajandream, M.A. and Barrell, B.G.

TITLE Direct Submission

JOURNAL Submitted (10-DEC-2000) Trypanosoma brucei genome sequencing project, Sanger Centre, The Wellcome Trust Genome Campus, Hinxton, Cambridge CB10 1SA. E-mail: barrell@sanger.ac.uk and nh@sanger.ac.uk

COMMENT Constructed at the Institute for Genomic Research (TIGR), Rockville, MD. Genomic DNA isolated from a cloned population of Trypanosoma brucei (FREG927/4 GPUT 10.1) was mechanically sheared to give a tight size distribution (4 kb). The v + i method used for the library construction is described in detail in Smith, R. and Venter, J.C. (Making small insert libraries for whole genome shotgun sequencing projects. In Genome Sequencing: A Practical Approach, eds. M. Vaudin and B. Barrell, Oxford University Press, 1999).

Email: nelsayed@tigr.org

Details of T. brucei sequencing at the Sanger Centre are available at http://www.sanger.ac.uk/Projects/T_brucei/.

Location/Qualifiers

1. 22

Source /organism="Trypanosoma brucei"

/strain="FREG927"

/db_xref="taxon:5691"

/clone="386h07"

BASE COUNT 0 a 4 c 0 g 18 t

ORIGIN

Query Match 0.6%; Score 16; DB 13; Length 22;

Best Local Similarity 100.0%; Pred. No. 1e+06;

Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaataaaaaa 2566

|||||

Db 20 GGAATAAAATAAAAAA 5

RESULT 22

AZ390689 23 bp DNA GSS 03-OCT-2000

LOCUS 1M0152A18F Mouse 10kb plasmid UUC1M library Mus musculus genomic

DEFINITION Clone UUC1M0152A18 F, DNA sequence.

ACCESSION AZ390689

VERSION AZ390689.1 GI:10505732

KEYWORDS GSS.

SOURCE house mouse.

ORGANISM Mus musculus

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.

REFERENCE 1 (bases 1 to 23)

AUTHORS Dunn, D., Aoyagi, A., Barber, M., Beacorn, T., Duval, B., Hamil, C., Islam, H., Longacre, S., Mahmoud, M., Meenen, E., Pedersen, F., Reilly, M., Rose, M., Rose, R., Stokes, R., Tingey, A., von Niederhausern, A. and Wright, D., Weiss, R.

TITLE Mouse whole genome scaffolding with paired end reads from 10kb plasmid inserts

JOURNAL Unpublished (2000)

COMMENT Contact: Robert B. Weiss

University of Utah

Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SIC, UT

84112, USA

Tel: 801 585 5606

Fax: 801 585 7177

Email: ddunn@genetics.utah.edu

Insert Length: 10000 Std Error: 0.00

Plate: 0152 row: A column: 18

Seq primer: CGTTGTAACGACGCCACT

Class: plasmid ends

High quality sequence stop: 23.

FEATURES

Source

1. 23

Location/Qualifiers

1. 23

/organism="Mus musculus"

/strain="C57BL/6J"

/db_xref="taxon:10090"

/clone="UUC1M0152A18"

/clone_11b="Mouse 10kb plasmid UUC1M library"

/sex="Male"

/lab_host="E. coli strain XL10-Gold, T1-resistant, F-"

/note="Vector: PWD42nv; Purified genomic DNA from M. musculus C57BL/6J (male) was obtained from the Jackson Laboratory Mouse DNA Resource

(<http://www.jax.org/resources/documents/dnares/>). The DNA was hydrodynamically sheared by repeated passage through a 0.005 inch orifice at constant velocity. The sheared DNA was blunt end-repaired with T4 DNA polymerase and T4 polynucleotide kinase. Adaptor oligonucleotides were ligated to the blunt ends in high molar excess. The adaptor DNA was purified and size-selected for a 9.5 to 10.5 kb range using preparative agarose gel electrophoresis. Vector DNA was prepared from a derivative of PWD42 (g1473211419b/AF129072.1), a copy-number inducible derivative of plasmid R1. The vector was ligated with adaptors complementary to the insert adaptors and purified. The sheared, adaptor mouse DNA was annealed to adaptor vector DNA, and transformed into chemically-competent E. coli XL10-Gold (Stratagene) cells and selected for ampicillin resistance."

BASE COUNT 16 a 0 c 7 g 0 t

ORIGIN

Query Match 0.6%; Score 16; DB 13; Length 23;

Best Local Similarity 100.0%; Pred. No. 1e+06;

Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaataaaaaa 2566

|||||

Db 8 GGAATAAAATAAAAAA 23

RESULT 23

AZ812579 24 bp DNA GSS 20-FEB-2001

LOCUS 2M0079A23F Mouse 10kb plasmid UUC1M library Mus musculus genomic

DEFINITION Clone UUC2M0079A23 F, DNA sequence.

ACCESSION AZ812579

VERSION AZ812579.1 GI:12981965

KEYWORDS GSS.

SOURCE house mouse.

ORGANISM Mus musculus

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.

REFERENCE 1 (bases 1 to 24)

AUTHORS Dunn, D., Aoyagi, A., Barber, M., Beacorn, T., Duval, B., Hamil, C., Islam, H., Longacre, S., Mahmoud, M., Meenen, E., Pedersen, F., Reilly, M., Rose, M., Rose, R., Stokes, R., Tingey, A., von Niederhausern, A. and Wright, D., Weiss, R.

TITLE Mouse whole genome scaffolding with paired end reads from 10kb plasmid inserts

JOURNAL Unpublished (2000)

COMMENT Contact: Robert B. Weiss

University of Utah

University of Utah

```

Ft      /db_xref="taxon:9606"
Ft      /organism="Homo sapiens"
Ft      /clone="DKFzp566J0846"

```


FT /clone_1lb="566 (synonym: hfkD2). Vector pAMP1; host
 FT X1-2bline: sites NotI + SalI"
 FT /dev_stage="fetal"
 FT /tissue_type="kidney"
 XX Sequence 25 BP; 1 A; 5 C; 0 G; 19 T; 0 other;

Query Match 0.6%; Score 16; DB 2; Length 25;
 Best Local Similarity 100.0%; Pred. No. 9.5e+05;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaataaaataa 2566
 |||||||
 Db 21 GGAATAAAATAAAATAA 6

RESULT 26
 HSM003450 standard; RNA; EST; 25 BP.
 ID HSM003450
 XX AL038974;
 XX AL038974.1

SV 12-MAR-1999 (Rel. 59, Created)
 DT 12-MAR-1999 (Rel. 59, Last updated, Version 1)
 XX Homo sapiens mRNA; EST DKFZp566a224_r1 (from clone DKFZp566a224)
 DE EST; expressed sequence tag.
 XX

OS Homo sapiens (human)
 OC Eukaryota; Metazoa; Chordata; Vertebrata; Euteleostomi; Mammalia;
 OC Eutheria; Primates; Catarrhini; Homiidae; Homo.
 XX

[1]
 1-25
 RA Boecker H., Boecker M., Brandt P., Mewes W., Gassenhuber J., Wiemann S.;
 RL Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
 RL MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
 XX

CC Clone from S. Wiemann, sequenced by GBF within the CDNA
 CC sequencing consortium of the German Genome Project
 CC No s1 sequence available at the RZPD in Berlin
 CC This clone is available at the RZPD in Berlin
 CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
 CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
 XX

FH Key Location/Qualifiers
 FH source 1.25
 FT /db_xref="taxon:9606"
 FT /organism="Homo sapiens"
 FT /clone="DKFZp566a224"

FT /clone_1lb="566 (synonym: hfkD2). Vector pAMP1; host
 FT X1-2bline: sites NotI + SalI"
 FT /dev_stage="fetal"
 FT /tissue_type="kidney"
 FT
 XX

SO Sequence 25 BP; 18 A; 0 C; 7 G; 0 T; 0 other;

Query Match 0.6%; Score 16; DB 2; Length 25;
 Best Local Similarity 100.0%; Pred. No. 9.5e+05;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaataaaataa 2566
 |||||||
 Db 6 GGAATAAAATAAAATAA 21

RESULT 27
 AZ832800/c GSS 20-FEB-2001
 LOCUS 2M0113M21F Mouse 10kb plasmid UUC1M library Mus musculus genomic
 DEFINITION clone UUC2M0113M21 F, DNA sequence.
 ACCESSION AZ832800
 VERSION 1
 KEYWORDS GSS.
 SOURCE house mouse.
 ORGANISM Mus musculus
 Eukaryota; Metazoa; Chordata; Vertebrata; Euteleostomi;
 Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
 1 (bases 1 to 25)
 REFERENCE Dunn,D., Aoyagi,A., Barber,M., Beacorn,T., Duval,B., Hamil,C.,
 Islam,H., Longacre,S., Mahmood,M., Meenen,E., Pedersen,T., Rellly
 M., Rose,M., Rose,R., Stokes,R., Tingey,A., von Niederhausen,A.
 and Wright,D., Weiss,R.
 Mouse whole genome scaffolding with paired end reads from 10kb
 plasmid inserts
 Unpublished (2000)
 CONTACT Robert B. Weiss
 University of Utah Genome Center
 University of Utah
 Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT
 84112, USA
 TEL: 801 585 5606
 FAX: 801 585 7177
 EMAIL: ddunn@genetics.utah.edu
 INSERT LENGTH: 10000 Std Error: 0.00
 PLATE: 0113 row: M column: 21
 SEQ PRIMER: CGTGTAAACGACGCCACGT
 CLASS: plasmid ends
 High quality sequence stop: 25.

FEATURES
 source 1.25
 Location/Qualifiers
 1.25
 /organism="Mus musculus"
 /strain="C57BL/6J"
 /db_xref="taxon:10090"
 /clone="UUC2M0113M21"
 /clone_1lb="Mouse 10kb plasmid UUC1M library"
 /sex="Male"
 /lab_host="E. Coli strain XL10-Gold, Tl-resistant, F-"
 /note="Vector: pMD42nv: Purified genomic DNA from M.
 musculus C57BL/6J (male) was obtained from the Jackson
 Laboratory Mouse DNA Resource
 (http://www.jax.org/resources/documents/dnares/). The DNA
 was hydrodynamically sheared by repeated passage through a
 0.005 inch orifice at constant velocity. The sheared DNA
 was blunt end-repaired with T4 DNA polymerase and T4
 polynucleotide kinase. Adaptor oligonucleotides were
 ligated to the blunt ends in high molar excess. The
 adaptor DNA was purified and size-selected for a 9.5 to
 10.5 kb range using preparative agarose gel
 electrophoresis. Vector DNA was prepared from a derivative
 of pMD2 (g11473211419b1AF129072.1) a copy-number
 inducible derivative of plasmid R1. The vector was ligated
 with adaptors complementary to the insert adaptors and
 purified. The sheared, adaptor mouse DNA was annealed to
 adaptor vector DNA, and transformed into
 chemically-competent E. coli XL10-Gold (Stratagene) cells
 and selected for ampicillin resistance."

BASE COUNT 0 a 2 c 0 g 23 t
 ORIGIN

Query Match 0.6%; Score 16; DB 13; Length 25;
 Best Local Similarity 100.0%; Pred. No. 9.5e+05;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaataaaataa 2566
 |||||||
 Db 25 GGAATAAAATAAAATAA 10

```

RESULT 28
AZ837511/c 25 bp DNA GSS 20-FEB-2001
LOCUS
DEFINITION
2M0132N17R Mouse 10kb plasmid UUGC1M library Mus musculus genomic
clone UUGC2M0132N17 R. DNA sequence.
ACCESSION
AZ837511
VERSION
AZ837511.1 GI:13007419
KEYWORDS
GSS.
SOURCE
house mouse.
ORGANISM
Mus musculus
REFERENCE
Mammalia; Metazoa; Chordata; Cranialata; Vertebrata; Euteleostomi;
Eukaryota; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
1 (bases 1 to 25)
Dunn, D., Aoyagi, A., Barber, M., Beacorn, T., Duval, B., Hamill, C.,
Islam, H., Longacre, S., Mahmoud, M., Meenen, E., Pedersen, T., Reilly,
M., Rose, M., Rose, R., Stokes, R., Tingey, A., von Niederhausern, A.
and Wright, D., Weiss, R.
Mouse whole genome scaffolding with paired end reads from 10kb
plasmid inserts
Unpublished (2000)
Contact: Robert B. Weiss:
University of Utah Genome Center
Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT
84112, USA
Tel: 801 585 5606
Fax: 801 585 7177
Email: ddunn@genetics.utah.edu
Insert length: 10000 Std Error: 0.00
Plate: 0132 row: N column: 17
Seq primer: CACACAGCAACACAGCTATGACC
Class: plasmid ends
High quality sequence stop: 25.
Location/Qualifiers
1..25
/organism="Mus musculus"
/strain="C57BL/6J"
/db_xref="taxon:10090"
/clone="UUGC2M0132N17"
/clone.lib="Mouse 10kb plasmid UUGC1M library"
/sex="Male"
/lab_host="E. coli strain XL10-Gold, T1-resistant, F-"
/notes="Vector: PWD42nv; Purified genomic DNA from M.
musculus C57BL/6J (male) was obtained from the Jackson
Laboratory Mouse DNA Resource
(http://www.jax.org/resources/documents/dnares/). The DNA
was hydrodynamically sheared by repeated passage through a
0.005 inch orifice at constant velocity. The sheared DNA
was blunt end-repaired with T4 DNA polymerase and T4
polynucleotide kinase. Adaptor oligonucleotides were
ligated to the blunt ends in high molar excess. The
adapted DNA was purified and size-selected for a 9.5 to
10.5 kb range using preparative agarose gel.
electrophoresis. Vector DNA was prepared from a derivative
of PWD42 (g11473211419b1AF129072.1), a copy-number
inducible derivative of plasmid RL. The vector was ligated
with adaptors complementary to the insert adaptors and
purified. The sheared, adapted mouse DNA was annealed to
adapted vector DNA, and transformed into
chemically-competent E. coli XL10-Gold (Stratagene) cells
and selected for ampicillin resistance."
BASE COUNT
0 a 10 c 0 g 15 t
ORIGIN
Query Match 0.6%; Score 16; DB 13; Length 25;
Best Local Similarity 100.0%; Pred. No. 9.5e+05;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 2551 ggaataaaaaa 2566
17 GGAATAAAAAA 2

```

```

RESULT 29
TA324F07P/c 25 bp DNA GSS 13-DEC-2000
LOCUS
DEFINITION
T. brucei sheared genomic DNA clone 324f07, forward sequence,
genomic survey sequence.
ACCESSION
AL493403
VERSION
AL493403.1 GI:11867768
KEYWORDS
GSS.
SOURCE
Trypanosoma brucei.
ORGANISM
Trypanosoma brucei
Eukaryota; Euglenozoa; Kinetoplastida; Trypanosomatidae;
Trypanosoma.
1 (bases 1 to 25)
Hall, N., Bowman, S., Lennard, N.J., Doggett, J., Atkin, R.,
Chillingworth, C., Ormond, D., Harris, B., El-Sayed, N., Hou, L.,
Melville, S.E., Rajadream, M.A. and Barrell, B.G.
Direct Submission
Submitted (10-DEC-2000) Trypanosoma brucei genome sequencing
project, Sanger Centre, The Wellcome Trust Genome Campus, Hinxton,
Cambridge CB10 1SA, E-mail: barrell@sanger.ac.uk and
nhs@sanger.ac.uk
Constructed at the Institute for Genomic Research (TIGR),
Rockville, MD. Genomic DNA isolated from a cloned population of
Trypanosoma brucei (TREU927/4 GUTat 10.1) was mechanically sheared
to give a tight size distribution (
4 kb). The v + i method used for the library construction is
described in detail in Smith, H. and Venter, J.C. (Making small
insert libraries for whole genome shotgun sequencing projects. In
Genome Sequencing: A Practical Approach, eds. M. Vaudin and B.
Barrell, Oxford University Press, 1999).
Email: nelsayed@tigr.org
Details of T. brucei sequencing at the Sanger Centre are available
at http://www.sanger.ac.uk/projects/T_brucei/.
Location/Qualifiers
1..25
/organism="Trypanosoma brucei"
/strain="TREU927"
/db_xref="taxon:5691"
/clone="324f07"
BASE COUNT
0 a 9 c 0 g 16 t
ORIGIN
Query Match 0.6%; Score 16; DB 13; Length 25;
Best Local Similarity 100.0%; Pred. No. 9.5e+05;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 2551 ggaataaaaaa 2566
18 GGAATAAAAAA 3

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```

RESULT 30
HSM001401
ID HSM001401 standard; RNA; EST; 26 BP.
XX
AC AL037076;
XX
SV AL037076.1;
XX
DT 12-MAR-1999 (Rel. 59, Created)
DT 12-MAR-1999 (Rel. 59, last updated, Version 1)
XX
DE Homo sapiens mRNA; EST DKFZp564K1764_r1 (from clone DKFZp564K1764)
XX
KW EST; expressed sequence tag.
XX
OS Homo sapiens (human)
XX
OC Eukaryota; Metazoa; Chordata; Cranialata; Vertebrata; Euteleostomi; Mammalia;
XX
OC Eutheria; Primates; Catarrhini; Homiinae; Homo.
XX

```

RN [1]
 RA 1-26
 RA Diesterhoeft A., Lauber J., Mewes W., Gassenhuber J., Wiemann S.;
 RT Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
 RL MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
 XX
 CC Clone from S. Wiemann, sequenced by Qiagen within the CDNA
 CC sequencing consortium of the German Genome Project
 CC No s1 sequence available
 CC This clone is available at the RZPD in Berlin
 CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
 CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
 XX
 FH Key Location/Qualifiers
 FT 1. 26
 FT source
 FT /db_xref="taxon:9606"
 FT /organism="Homo sapiens"
 FT /clone="DKFZp564K1764"
 FT /clone_1lb="564 (synonym: hfbz2). Vector pAMP1; host
 FT X1-2blue; sites NotI + SalI"
 FT /dev_stage="fetal"
 FT /tissue_type="brain"
 XX
 SO Sequence 26 BP; 18 A; 0 C; 8 G; 0 T; 0 other;

Query Match 0.6%; Score 16; DB 2; Length 26;
 Best Local Similarity 100.0%; Pred. No. 9.2e+05;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaataaaataaa 2566
 Db 7 GGAATAAAATAAAATAAA 22

RESULT 31
 ID HSM001421 standard; RNA; EST; 26 BP.
 XX
 AC AL037096;
 XX
 SV AL037096.1
 XX
 DT 12-MAR-1999 (Rel. 59, Created)
 DT 12-MAR-1999 (Rel. 59, Last updated, Version 1)
 XX
 DE Homo sapiens mRNA; EST DKFZp564L2164_r1 (from clone DKFZp564L2164)
 XX
 KW EST; expressed sequence tag.
 XX
 OS Homo sapiens (human)
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
 OC Eutheria; Primates; Catarrhini; Homiidae; Homo.
 XX
 RN [1]
 RN 1-26
 RA Diesterhoeft A., Lauber J., Mewes W., Gassenhuber J., Wiemann S.;
 RT Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
 RL MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
 XX
 CC Clone from S. Wiemann, sequenced by Qiagen within the CDNA
 CC sequencing consortium of the German Genome Project
 CC No s1 sequence available
 CC This clone is available at the RZPD in Berlin
 CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
 CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
 XX
 FH Key Location/Qualifiers
 FT 1. 26
 FT source

FT /db_xref="taxon:9606"
 FT /organism="Homo sapiens"
 FT /clone="DKFZp564L2164"
 FT /clone_1lb="564 (synonym: hfbz2). Vector pAMP1; host
 FT X1-2blue; sites NotI + SalI"
 FT /dev_stage="fetal"
 FT /tissue_type="brain"
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Query Match 0.6%; Score 16; DB 2; Length 26;
 Best Local Similarity 100.0%; Pred. No. 9.2e+05;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaataaaataaa 2566
 Db 4 GGAATAAAATAAAATAAA 19

RESULT 32
 ID HSM002179 standard; RNA; EST; 26 BP.
 XX
 AC AL037846;
 XX
 SV AL037846.1
 XX
 DT 12-MAR-1999 (Rel. 59, Created)
 DT 12-MAR-1999 (Rel. 59, Last updated, Version 1)
 XX
 DE Homo sapiens mRNA; EST DKFZp564I177_r1 (from clone DKFZp564I177)
 XX
 KW EST; expressed sequence tag.
 XX
 OS Homo sapiens (human)
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
 OC Eutheria; Primates; Catarrhini; Homiidae; Homo.
 XX
 RN [1]
 RN 1-26
 RA Bloeker H., Boecher M., Brandt P., Mewes W., Gassenhuber J., Wiemann S.;
 RT Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
 RL MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
 XX
 CC Clone from S. Wiemann, sequenced by GBF within the CDNA
 CC sequencing consortium of the German Genome Project
 CC No s1 sequence available
 CC This clone is available at the RZPD in Berlin
 CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
 CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
 XX
 FH Key Location/Qualifiers
 FT 1. 26
 FT source
 FT /db_xref="taxon:9606"
 FT /organism="Homo sapiens"
 FT /clone="DKFZp564I177"
 FT /clone_1lb="564 (synonym: hfbz2). Vector pAMP1; host
 FT X1-2blue; sites NotI + SalI"
 FT /dev_stage="fetal"
 FT /tissue_type="brain"
 XX
 SO Sequence 26 BP; 18 A; 0 C; 8 G; 0 T; 0 other;

Query Match 0.6%; Score 16; DB 2; Length 26;
 Best Local Similarity 100.0%; Pred. No. 9.2e+05;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaataaaataaa 2566
 Db 7 GGAATAAAATAAAATAAA 22

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Db      7 GGAAGAAAAA 22
RESULT 33
ID      HSM002201/c standard; RNA; EST; 26 BP.
XX      AL037868;
XX      AL037868.1
SV      12-MAR-1999 (Rel. 59, Created)
DT      12-MAR-1999 (Rel. 59, Last updated, Version 1)
XX      Homo sapiens mRNA; EST DKFZp564J177_s1 (from clone DKFZp564J177)
XX      EST; expressed sequence tag.
XX      Homo sapiens (human)
OS      Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
OC      Eutheria; Primates; Catarrhini; Homnidae; Homo.
XX      [1]
XX      Bloecker H., Boecker M., Brandt P., Mewes W., Gassenhuber J., Wiemann S.;
RP      1-26
RA      Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
RT      MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
RL      MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
XX      Clone from S. Wiemann, sequenced by GBF within the CDNA
CC      sequencing consortium of the German Genome Project
CC      r1 sequence also available
CC      This clone is available at the RZPD in Berlin
CC      Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
CC      Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
XX      Key      Location/Qualifiers
FH      source      1..26
FT      /db_xref="taxon:9606"
FT      /organism="Homo sapiens"
FT      /clone="DKFZp564J177"
FT      /clone_11b="564 (synonym: hfb2). Vector pAMP1; host
FT      X1-2blue; sites: NotI + SalI"
FT      /dev_stage="fetal"
FT      /tissue_type="brain"
XX      Sequence 26 BP; 0 A; 8 C; 0 G; 18 T; 0 other;
SQ
Query Match      0.6%; Score 16; DB 2; Length 26;
Best Local Similarity 100.0%; Pred. No. 9.2e+05;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
OY      2551 ggaagaaaaa 2566
DB      20 GGAAGAAAAA 5
RESULT 34
ID      HSM003490 standard; RNA; EST; 26 BP.
XX      AL039014;
XX      AL039014.1
SV      12-MAR-1999 (Rel. 59, Created)
DT      12-MAR-1999 (Rel. 59, Last updated, Version 1)
XX      Homo sapiens mRNA; EST DKFZp566C074_r1 (from clone DKFZp566C074)
XX      EST; expressed sequence tag.

```

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XX      Homo sapiens (human)
OS      Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
OC      Eutheria; Primates; Catarrhini; Homnidae; Homo.
XX      [1]
XX      Bloecker H., Boecker M., Brandt P., Mewes W., Gassenhuber J., Wiemann S.;
RP      1-26
RA      Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
RT      MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
RL      MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
XX      Clone from S. Wiemann, sequenced by GBF within the CDNA
CC      sequencing consortium of the German Genome Project
CC      No s1 sequence available
CC      This clone is available at the RZPD in Berlin
CC      Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
CC      Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
XX      Key      Location/Qualifiers
FH      source      1..26
FT      /db_xref="taxon:9606"
FT      /organism="Homo sapiens"
FT      /clone="DKFZp566C074"
FT      /clone_11b="566 (synonym: hfb2). Vector pAMP1; host
FT      X1-2blue; sites: NotI + SalI"
FT      /dev_stage="fetal"
FT      /tissue_type="kidney"
XX      Sequence 26 BP; 18 A; 0 C; 8 G; 0 T; 0 other;
SQ
Query Match      0.6%; Score 16; DB 2; Length 26;
Best Local Similarity 100.0%; Pred. No. 9.2e+05;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
OY      2551 ggaagaaaaa 2566
DB      7 GGAAGAAAAA 22

```

```

RESULT 35
ID      HSM003529 standard; RNA; EST; 26 BP.
XX      AL039053;
XX      AL039053.1
SV      12-MAR-1999 (Rel. 59, Created)
DT      12-MAR-1999 (Rel. 59, Last updated, Version 1)
XX      Homo sapiens mRNA; EST DKFZp566F04_r1 (from clone DKFZp566F04)
XX      EST; expressed sequence tag.
XX      Homo sapiens (human)
OS      Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
OC      Eutheria; Primates; Catarrhini; Homnidae; Homo.
XX      [1]
XX      Bloecker H., Boecker M., Brandt P., Mewes W., Gassenhuber J., Wiemann S.;
RP      1-26
RA      Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
RT      MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
RL      MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY
XX      Clone from S. Wiemann, sequenced by GBF within the CDNA
CC      sequencing consortium of the German Genome Project
CC      No s1 sequence available
CC      This clone is available at the RZPD in Berlin
CC      Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059

```

```

CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
XX
FH Key Location/Qualifiers
FH source
FT 1.26
FT /db_xref="taxon:9606"
FT /organism="Homo sapiens"
FT /clone="DKFZp566J14"
FT /clone_lib="566 (synonym: hfk2). Vector pAMP1; host
FT X1-2blue; sites NotI + SalI"
FT /dev_stage="fetal"
FT /tissue_type="kidney"
XX
XX Sequence 26 BP; 18 A; 0 C; 8 G; 0 T; 0 other;

Query Match
Best Local Similarity 100.0%; Score 16; DB 2; Length 26;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaagaaagaaagaaagaa 2566
Db 7 GGAAGAAAGAAAGAAAGAA 22

RESULT 36
HSM003586
ID HSM003586 standard; RNA; EST; 26 BP.
XX
XX AL039110;
XX
XX AL039110.1
XX
XX 12-MAR-1999 (Rel. 59, Created)
XX 12-MAR-1999 (Rel. 59, Last updated, Version 1)
XX Homo sapiens mRNA; EST DKFZp566J14_r1 (from clone DKFZp566J14)
XX
XX EST; expressed sequence tag.
XX
XX Homo sapiens (human)
XX Eukaryota; Metazoa; Chordata; Vertebrata; Euteleostomi; Mammalia;
XX Eutheria; Primates; Catarrhini; Homiidae; Homo.
XX
XX [1]
XX 1.26
XX Bloeker H., Boecker M., Brandt P., Mewes W., Gassenhuber J., Wiemann S.;
XX Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
XX MIPS, Am Klopferplatz 18a D-82152 Martinsried, GERMANY
XX
XX Clone from S. Wiemann, sequenced by GBF within the CDNA
XX sequencing consortium of the German Genome Project
XX No s1 sequence available
XX This clone is available at the RZPD in Berlin
XX Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
XX Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
XX
XX Key Location/Qualifiers
XX
XX source
XX 1.26
XX /db_xref="taxon:9606"
XX /organism="Homo sapiens"
XX /clone="DKFZp566J14"
XX /clone_lib="566 (synonym: hfk2). Vector pAMP1; host
XX X1-2blue; sites NotI + SalI"
XX /dev_stage="fetal"
XX /tissue_type="kidney"
XX
XX Sequence 26 BP; 19 A; 0 C; 7 G; 0 T; 0 other;

Query Match
0.6%; Score 16; DB 2; Length 26;

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Best Local Similarity 100.0%; Pred. No. 9.2e+05;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaagaaagaaagaaagaa 2566
Db 6 GGAAGAAAGAAAGAAAGAA 21

RESULT 37
HSM001542/C
ID HSM001542 standard; RNA; EST; 27 BP.
XX
XX AL037217;
XX
XX AL037217.1
XX
XX 12-MAR-1999 (Rel. 59, Created)
XX 12-MAR-1999 (Rel. 59, Last updated, Version 1)
XX Homo sapiens mRNA; EST DKFZp564B069_s1 (from clone DKFZp564B069)
XX
XX EST; expressed sequence tag.
XX
XX Homo sapiens (human)
XX Eukaryota; Metazoa; Chordata; Vertebrata; Euteleostomi; Mammalia;
XX Eutheria; Primates; Catarrhini; Homiidae; Homo.
XX
XX [1]
XX 1-27
XX Ansoerge W., Winkner U., Mewes W., Gassenhuber J., Wiemann S.;
XX Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.
XX MIPS, Am Klopferplatz 18a D-82152 Martinsried, GERMANY
XX
XX Clone from S. Wiemann, sequenced by EMBL within the CDNA
XX sequencing consortium of the German Genome Project
XX r1 sequence also available
XX This clone is available at the RZPD in Berlin
XX Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
XX Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
XX
XX Key Location/Qualifiers
XX
XX source
XX 1.27
XX /db_xref="taxon:9606"
XX /organism="Homo sapiens"
XX /clone="DKFZp564B069"
XX /clone_lib="564 (synonym: hfk2). Vector pAMP1; host
XX X1-2blue; sites NotI + SalI"
XX /dev_stage="fetal"
XX /tissue_type="brain"
XX
XX Sequence 27 BP; 0 A; 8 C; 0 G; 19 T; 0 other;

Query Match
0.6%; Score 16; DB 2; Length 27;
Best Local Similarity 100.0%; Pred. No. 9e+05;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaagaaagaaagaaagaa 2566
Db 21 GGAAGAAAGAAAGAAAGAA 6

RESULT 38
HSM002015
ID HSM002015 standard; RNA; EST; 27 BP.
XX
XX AL037684;
XX
XX AL037684.1
XX
XX 12-MAR-1999 (Rel. 59, Created)

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XX Sequence 27 BP; 0 A; 9 C; 0 G; 18 T; 0 other;

Query Match 0.6%; Score 16; DB 2; Length 27;
Best Local Similarity 100.0%; Pred. No. 9e+05;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaataaaataa 2566
|||||
DB 20 GGAATAAAATAAAATAA 5

RESULT 41
HSM003588/c
ID HSM003588 standard; RNA; EST; 27 BP.

AL039112.1

SV 12-MAR-1999 (Rel. 59, Created)
XX 12-MAR-1999 (Rel. 59, Last updated, Version 1)

DE Homo sapiens mRNA; EST DKFZp566J184_s1 (from clone DKFZp566J184)

XX EST: expressed sequence tag.

OS Homo sapiens (human)
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
XX Eutheria; Primates; Catarrhini; Homiidae; Homo.

[1]

RA Bloeker H., Boecker M., Brandt P., Mewes W., Gassenhuber J., Wiemann S.;

RT Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.

RL MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY

XX Clone from S. Wiemann, sequenced by GBF within the CDNA

CC sequencing consortium of the German Genome Project

CC No r1 sequence available
CC This clone is available at the RZPD in Berlin

CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059

CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de

XX Key Location/Qualifiers

FT source 1. 27
FT /db_xref="taxon:9606"
FT /organism="Homo sapiens"
FT /clone="DKFZp566J184"
FT /clone_11b="566 (synonym: hfkcd2). Vector pAMP1; host
FT X1-2blue; sites NotI + SalI"
FT /dev_stage="fetal"
FT /tissue_type="kidney"
XX Sequence 27 BP; 0 A; 8 C; 0 G; 19 T; 0 other;

Query Match 0.6%; Score 16; DB 2; Length 27;
Best Local Similarity 100.0%; Pred. No. 9e+05;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaataaaataa 2566
|||||
DB 21 GGAATAAAATAAAATAA 6

RESULT 42
HSM003597
ID HSM003597 standard; RNA; EST; 27 BP.

AC AL039121;

XX AL039121.1

SV 12-MAR-1999 (Rel. 59, Created)

DT 12-MAR-1999 (Rel. 59, Last updated, Version 1)

DE Homo sapiens mRNA; EST DKFZp566J084_r1 (from clone DKFZp566J084)

XX EST: expressed sequence tag.

OS Homo sapiens (human)
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia;
XX Eutheria; Primates; Catarrhini; Homiidae; Homo.

[1]

RA Bloeker H., Boecker M., Brandt P., Mewes W., Gassenhuber J., Wiemann S.;

RT Submitted (12-MAR-1999) to the EMBL/GenBank/DBJ databases.

RL MIPS, Am Klopferspitz 18a D-82152 Martinsried, GERMANY

XX Clone from S. Wiemann, sequenced by GBF within the CDNA

CC sequencing consortium of the German Genome Project

CC No s1 sequence available
CC This clone is available at the RZPD in Berlin

CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059

CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de

XX Key Location/Qualifiers

FT source 1. 27
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FT /organism="Homo sapiens"
FT /clone="DKFZp566J084"
FT /clone_11b="566 (synonym: hfkcd2). Vector pAMP1; host
FT X1-2blue; sites NotI + SalI"
FT /dev_stage="fetal"
FT /tissue_type="kidney"
XX Sequence 27 BP; 19 A; 0 C; 8 G; 0 T; 0 other;

Query Match 0.6%; Score 16; DB 2; Length 27;
Best Local Similarity 100.0%; Pred. No. 9e+05;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaataaaataaaataa 2566
|||||
DB 7 GGAATAAAATAAAATAA 22

RESULT 43

AV741507 27 bp mRNA EST 17-OCT-2000

LOCUS AV741507 CB Homo sapiens cDNA clone CEMAC05 5', mRNA sequence.

DEFINITION AV741507

ACCESSION AV741507.1 GI:10859088

VERSION AV741507.1

KEYWORDS EST.

SOURCE human.

ORGANISM Homo sapiens

REFERENCE 1 (bases 1 to 27)

AUTHORS Zhang, Q., Ye, M., Wu, X., Gu, J., Huang, Q., Zhou, J., Shen, Y., Han, Z.,

Chen, S., Mao, M., and Chen, Z.

TITLE Homo sapiens CB library cDNA clones

JOURNAL Unpublished (2000)

COMMENT Contact: Zhu Chen

Shanghai Institute of Hematology, Rui-Jin Hospital

197 Rui-Jin II Road, Shanghai 200025, P. R. China

Tel: 86-21-64740490

Fax: 86-21-64743206

Email: mbshiens.stn.sh.cn
This clone is available at Shanghai Hematology Institute in Shanghai.

Chinese National Human Genome Center at Shanghai
351 Guo Shoujing Road, Zhangjiang Hi-Tech Park, Pudong.

FEATURES

source

1.27

/organism="Homo sapiens"

/db_xref="taxon:9606"

/clone="CBMAMC05"

/clone.lib="CB"

/tissue.type="cord blood"

/cell_type="CD34+ hematopoietic stem/progenitor cell"

/lab_host="BM25.8"

/note="Vector: pBluescript; Site:1: EcoRI; The insert is cloned randomly with the EcoRI digestion"

BASE COUNT

17 a 2 c 6 g 0 t 2 others

ORIGIN

Query Match 0.6%; Score 16; DB 10; Length 27;

Best Local Similarity 100.0%; Pred. No. 9e+05; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaataaaataaa 2566

Db 11 GGAAAAAAAAAAAAA 26

RESULT 44

N29432/c

LOCUS

DEFINITION N29432 27 bp mRNA EST 05-JAN-1996

YW86110.s1 Soares,Placenta,8to9weeks,2NBHP8to9M Homo sapiens CDNA

clone IMAGE:259171 3' similar to gb:X64559 TETRAPECTIN PRECURSOR

(HUMAN);, mRNA sequence.

ACCESSION N29432

VERSION N29432.1

KEYWORDS

SOURCE

ORGANISM

human.

Homo sapiens

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Primates; Catarrhini; Homnidae; Homo.

REFERENCE

AUTHORS

Hillier,L., Clark,N., Dubuque,T., Elliston,K., Hawkins,M., Holman

M., Hultman,M., Kucaba,T., Le,M., Lennon,G., Marris,M., Parsons,J.,

Rifkin,L., Rohlfing,T., Soares,M., Tan,F., Trevaskis,E., Waterston

R., Williamson,A., Wohlmann,P. and Wilson,R.

The WashU-Merck EST Project

Unpublished (1995)

Contact: Wilson RK

Washington University School of Medicine

4444 Forest Park Parkway, Box 8501, St. Louis, MO 63108

Tel: 314 286 1800

Fax: 314 286 1810

Email: estewatson.wustl.edu

High quality sequence starts: 1

High quality sequence stops: 1

Source: IMAGE Consortium; LNLN

This clone is available royalty-free through LNLN; contact the

IMAGE Consortium (info@image.lnl.gov) for further information.

Trace considered overall poor quality

Seq primer: m13 -40 forward

High quality sequence stop: 1.

FEATURES

source

1.27

/organism="Homo sapiens"

/db_xref="GDB:3888877"

/db_xref="taxon:9606"

/clone="IMAGE:259171"

/clone.lib="Soares,Placenta,8to9weeks,2NBHP8to9M"

/dev_stage="two placenta: one from 8 weeks and another

from 9 weeks post conception"

/lab_host="DHI0B (ampicillin resistant)"

BASE COUNT

0 a 3 c 0 g 24 t

ORIGIN

Query Match 0.6%; Score 16; DB 11; Length 27;

Best Local Similarity 100.0%; Pred. No. 9e+05; Mismatches 0; Indels 0; Gaps 0;

QY 2551 ggaataaaataaaataaa 2566

Db 25 GGAAAAAAAAAAAAA 10

RESULT 45

R31539/c

LOCUS

DEFINITION R31539 27 bp mRNA EST 28-APR-1995

YH72605.s1 Soares,Placenta,Nb2HP Homo sapiens CDNA clone

IMAGE:135296 3' similar to gb:X53463 GLUTATHIONE

PEROXIDASE-GASTROINTESTINAL (HUMAN);, mRNA sequence.

ACCESSION R31539

VERSION R31539.1

KEYWORDS

SOURCE

ORGANISM

human.

Homo sapiens

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Primates; Catarrhini; Homnidae; Homo.

REFERENCE

AUTHORS

Hillier,L., Clark,N., Dubuque,T., Elliston,K., Hawkins,M., Holman

M., Hultman,M., Kucaba,T., Le,M., Lennon,G., Marris,M., Parsons,J.,

Rifkin,L., Rohlfing,T., Soares,M., Tan,F., Trevaskis,E., Waterston

R., Williamson,A., Wohlmann,P. and Wilson,R.

The WashU-Merck EST Project

Unpublished (1995)

Contact: Wilson RK

Washington University School of Medicine

4444 Forest Park Parkway, Box 8501, St. Louis, MO 63108

Tel: 314 286 1800

Fax: 314 286 1810

Email: estewatson.wustl.edu

High quality sequence starts: 1

High quality sequence stops: 1

Source: IMAGE Consortium; LNLN

This clone is available royalty-free through LNLN; contact the

IMAGE Consortium (info@image.lnl.gov) for further information.

Trace considered overall poor quality

Insert Length: 367 Std Error: 0.00

Seq primer: -21m13

High quality sequence stop: 1.

FEATURES

source

1.27

/organism="Homo sapiens"

/db_xref="GDB:541217"

/db_xref="taxon:9606"

/clone="IMAGE:135296"

/clone.lib="Soares,Placenta,Nb2HP"

/sex="Female"

/dev_stage="Placenta obtained at birth (full term)"

/lab_host="DHI0B (ampicillin resistant)"

/note="Organ: placenta; Vector: pT73D (Pharmacia) with a

modified polylinker; Site:1: Not I; Site:2: Eco RI; 1st

strand cDNA was primed with Not I and Eco RI sites of a modified pT73 vector

(Pharmacia). Library constructed by Bento Soares and

M.Fatima Bernaldo."

double-stranded cDNA was ligated to Eco RI adaptors

GenCore version 4.5
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OM nucleic - nucleic search, using sw model

Run on: April 20, 2002, 06:44:33 ; Search time 3629.2 Seconds

(without alignments)
11664.213 Million cell updates/sec

Title: US-10-024-396-3

Sequence: 1 cytcgcgcgtccgcgtctcct.....aaatgaaaaa 2566

Scoring table:

Gapop 60.0 , Gapext 60.0

Searched: 1472140 seqs, 8248589755 residues

Word size : 0

Total number of hits satisfying chosen parameters: 541028

Minimum DB seq length: 0

Maximum DB seq length: 50

Post-processing: Listing first 45 summaries

Database :

GenBank:
1: gb_ba:*
2: gb_htg:*
3: gb_in:*
4: gb_om:*
5: gb_ov:*
6: gb_pat:*
7: gb_ph:*
8: gb_pl:*
9: gb_pr:*
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13: gb_un:*
14: gb_vi:*
15: em_ba:*
16: em_fun:*
17: em_hum:*
18: em_in:*
19: em_om:*
20: em_or:*
21: em_ov:*
22: em_pat:*
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24: em_pl:*
25: em_ro:*
26: em_sts:*
27: em_sy:*
28: em_un:*
29: em_vi:*
30: em_htgo_hum:*
31: em_htgo_inv:*
32: em_htgo_rod:*
33: em_htg_hum:*
34: em_htg_inv:*
35: em_htg_rod:*
36: em_htg_other:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB	ID	Description
1	31	1.2	34	6	AR112204	AR112204 Sequence
2	31	1.2	34	6	AR149246	AR149246 Sequence
3	26	1.0	33	6	AR096476	AR096476 Sequence
4	24	0.9	24	6	AR092018	AR092018 Sequence
5	24	0.9	24	6	AR112153	AR112153 Sequence
6	24	0.9	24	6	AR149195	AR149195 Sequence
7	23	0.9	23	6	AR112201	AR112201 Sequence
8	23	0.9	23	6	AR149243	AR149243 Sequence
9	23	0.9	31	6	AR092048	AR092048 Sequence
10	23	0.9	31	6	AR092050	AR092050 Sequence
11	23	0.9	31	6	AR112183	AR112183 Sequence
12	23	0.9	31	6	AR112185	AR112185 Sequence
13	23	0.9	31	6	AR149225	AR149225 Sequence
14	23	0.9	31	6	AR149227	AR149227 Sequence
15	18	0.7	36	6	AR096475	AR096475 Sequence
16	18	0.7	50	6	AX161232	AX161232 Sequence
17	17	0.7	20	6	AR092047	AR092047 Sequence
18	17	0.7	20	6	AR092049	AR092049 Sequence
19	17	0.7	20	6	AR112182	AR112182 Sequence
20	17	0.7	20	6	AR112184	AR112184 Sequence
21	17	0.7	20	6	AR149224	AR149224 Sequence
22	17	0.7	20	6	AR149226	AR149226 Sequence
23	17	0.7	25	6	AX043026	AX043026 Sequence
24	17	0.7	47	6	AR142905	AR142905 Sequence
25	16	0.6	18	6	E32461	E32461 Mammal-derl
26	16	0.6	20	6	E29883	E29883 HIV cofactor
27	16	0.6	22	6	AR142908	AR142908 Sequence
28	16	0.6	23	6	AX115478	AX115478 Sequence
29	16	0.6	25	6	A90999	A90999 Sequence 1
30	16	0.6	25	6	A91901	A91901 Sequence 5
31	16	0.6	25	6	AR106367	AR106367 Sequence
32	16	0.6	25	6	AR148371	AR148371 Sequence
33	16	0.6	25	6	AX032404	AX032404 Sequence
34	16	0.6	25	6	AX042532	AX042532 Sequence
35	16	0.6	25	6	AX042542	AX042542 Sequence
36	16	0.6	25	6	AX042571	AX042571 Sequence
37	16	0.6	25	6	AX042589	AX042589 Sequence
38	16	0.6	25	6	AX042627	AX042627 Sequence
39	16	0.6	25	6	AX042759	AX042759 Sequence
40	16	0.6	25	6	AX042823	AX042823 Sequence
41	16	0.6	25	6	AX042825	AX042825 Sequence
42	16	0.6	25	6	AX042827	AX042827 Sequence
43	16	0.6	25	6	AX042871	AX042871 Sequence
44	16	0.6	25	6	AX042878	AX042878 Sequence
45	16	0.6	25	13	AX032410	AX032410 Sequence

ALIGNMENTS

RESULT	1	PAT	16-MAY-2001
LOCUS	AR112204		
DEFINITION	Sequence 93 from patent US 6130041.		
ACCESSION	AR112204		
VERSION	AR112204.1 GI:14092104		
KEYWORDS	Unknown.		
SOURCE	Unknown.		
ORGANISM	Unclassified.		
REFERENCE	1 (bases 1 to 34)		
AUTHORS	Acton, S. Laurence.		
TITLE	Human intronic and polymorphic SR-BI nucleic acids and uses therefor		
JOURNAL	Patent: US 6130041-A 93 10-OCT-2000;		
FEATURES	Location/Qualifiers		
BASE COUNT	4 a 15 c 3 g 12 t		
ORIGIN	/organism="unknown"		

Query Match 1.2%; Score 31; DB 6; Length 34;
Best Local Similarity 100.0%; Pred. No. 4.1e-05;
Matches 31; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1085 cctgttctctccatcctcactcctca 1115
Db 1 CCTGTCTCTCTCCATCTCCTCTCA 31

RESULT 2
LOCUS AR149246 34 bp DNA PAT 08-AUG-2001
DEFINITION Sequence 93 from patent US 6228581.
ACCESSION AR149246
VERSION AR149246.1 GI:15113837
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE 1 (bases 1 to 34)
AUTHORS Acton,S.L. and Ordovas,J.M.
TITLE Human intronic and polymorphic SR-BI nucleic acids and uses therefor
JOURNAL Patent: US 6228581-A 93 08-MAY-2001;
FEATURES Location/Qualifiers
1..34
source /organism="unknown"

BASE COUNT 4 a 15 c 3 g 12 t
ORIGIN

Query Match 1.2%; Score 31; DB 6; Length 34;
Best Local Similarity 100.0%; Pred. No. 4.1e-05;
Matches 31; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1085 cctgttctctccatcctcactcctca 1115
Db 1 CCTGTCTCTCTCCATCTCCTCTCA 31

RESULT 3
LOCUS AR096476 33 bp DNA PAT 08-SEP-2000
DEFINITION Sequence 5 from patent US 6008014.
ACCESSION AR096476
VERSION AR096476.1 GI:10025312
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE 1 (bases 1 to 33)
AUTHORS Gimeno,C.J. and Acton,S.
TITLE Method of making lipid metabolic pathway compositions
JOURNAL Patent: US 6008014-A 5 28-DEC-1999;
FEATURES Location/Qualifiers
1..33
source /organism="unknown"

BASE COUNT 6 a 10 c 7 g 10 t
ORIGIN

Query Match 1.0%; Score 26; DB 6; Length 33;
Best Local Similarity 100.0%; Pred. No. 0.024;
Matches 26; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1576 gtgctgcaggaagaactgttagg 1601
Db 33 GTGCTGCAGGAAGCAAACTAGGCG 8

RESULT 4

AR092018/c AR092018 24 bp DNA PAT 08-SEP-2000
LOCUS AR149195/c
DEFINITION Sequence 42 from patent US 5998141.
ACCESSION AR092018
VERSION AR092018.1 GI:10018772
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE 1 (bases 1 to 24)
AUTHORS Acton,S.L. and Ordovas,J.M.
TITLE Human intronic and polymorphic SR-BI nucleic acids and uses therefor
JOURNAL Patent: US 5998141-A 42 07-DEC-1999;
FEATURES Location/Qualifiers
1..24
source /organism="unknown"

BASE COUNT 3 a 8 c 8 g 5 t
ORIGIN

Query Match 0.9%; Score 24; DB 6; Length 24;
Best Local Similarity 100.0%; Pred. No. 0.31;
Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 67 gacatggctgctccgccaagcg 90
Db 24 GACATGGCTGCTCCGCCAAGCG 1

RESULT 5
LOCUS AR112153 24 bp DNA PAT 16-MAY-2001
DEFINITION Sequence 42 from patent US 6130041.
ACCESSION AR112153
VERSION AR112153.1 GI:14092053
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE 1 (bases 1 to 24)
AUTHORS Acton,S.L. and Ordovas,J.M.
TITLE Human intronic and polymorphic SR-BI nucleic acids and uses therefor
JOURNAL Patent: US 6130041-A 42 10-OCT-2000;
FEATURES Location/Qualifiers
1..24
source /organism="unknown"

BASE COUNT 3 a 8 c 8 g 5 t
ORIGIN

Query Match 0.9%; Score 24; DB 6; Length 24;
Best Local Similarity 100.0%; Pred. No. 0.31;
Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 67 gacatggctgctccgccaagcg 90
Db 24 GACATGGCTGCTCCGCCAAGCG 1

RESULT 6
LOCUS AR149195 24 bp DNA PAT 08-AUG-2001
DEFINITION Sequence 42 from patent US 6228581.
ACCESSION AR149195
VERSION AR149195.1 GI:15113786
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE 1 (bases 1 to 24)
AUTHORS Acton,S.L. and Ordovas,J.M.
TITLE Human intronic and polymorphic SR-BI nucleic acids and uses therefor

BASE COUNT 3 a 8 c 8 g 5 t
ORIGIN

therefor
JOURNAL Patent: US 6228581-A 42 08-MAY-2001;
FEATURES Location/Qualifiers
Source 1..24
/organism="unknown"
BASE COUNT 3 a 8 c 8 g 5 t
ORIGIN

Query Match 0.9%; Score 24; DB 6; Length 24;
Best Local Similarity 100.0%; Pred. No. 0.31;
Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 67 gacatggctgctccgcaagcg 90
|||||
Db 24 GACATGGCTCTCTCCGCAAGCG 1

RESULT 7
AR112201/c. AR112201 23 bp DNA PAT 16-MAY-2001
LOCUS
DEFINITION Sequence 90 from patent US 6130041.
ACCESSION AR112201
VERSION AR112201.1 GI:14092101
KEYWORDS
SOURCE Unknown.
ORGANISM
REFERENCE Unclassified.
AUTHORS 1 (bases 1 to 23)
TITLE Acton,S.Laurene.
Human intronic and polymorphic SR-BI nucleic acids and uses therefor

JOURNAL Patent: US 6130041-A 90 10-OCT-2000;
FEATURES Location/Qualifiers
Source 1..23
/organism="unknown"

BASE COUNT 6 a 10 c 6 g 1 t
ORIGIN

Query Match 0.9%; Score 23; DB 6; Length 23;
Best Local Similarity 100.0%; Pred. No. 1.1;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 123 gctactgtgcgtgctgagcg 145
|||||
Db 23 GCTACTGTGCGCTGTGCTGGCG 1

RESULT 8
AR149243/c. AR149243 23 bp DNA PAT 08-AUG-2001
LOCUS
DEFINITION Sequence 90 from patent US 6228581.
ACCESSION AR149243
VERSION AR149243.1 GI:15113834
KEYWORDS
SOURCE Unknown.
ORGANISM
REFERENCE Unclassified.
AUTHORS 1 (bases 1 to 23)
TITLE Acton,S.L. and Ordovas,J.M.
Human intronic and polymorphic SR-BI nucleic acids and uses therefor

JOURNAL Patent: US 6228581-A 90 08-MAY-2001;
FEATURES Location/Qualifiers
Source 1..23
/organism="unknown"

BASE COUNT 6 a 10 c 6 g 1 t
ORIGIN

Query Match 0.9%; Score 23; DB 6; Length 23;
Best Local Similarity 100.0%; Pred. No. 1.1;

Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 123 gctactgtgcgtgctgagcg 145
|||||
Db 23 GCTACTGTGCGCTGTGCTGGCG 1

RESULT 9
AR092048/c. AR092048 31 bp DNA PAT 08-SEP-2000
LOCUS
DEFINITION Sequence 72 from patent US 5998141.
ACCESSION AR092048
VERSION AR092048.1 GI:10018802
KEYWORDS
SOURCE Unknown.
ORGANISM
REFERENCE Unclassified.
AUTHORS 1 (bases 1 to 31)
TITLE Acton,S.Laurene.
Intronic and polymorphic SR-BI nucleic acids and uses therefor

JOURNAL Patent: US 5998141-A 72 07-DEC-1999;
FEATURES Location/Qualifiers
Source 1..31
/organism="unknown"

BASE COUNT 7 a 6 c 12 g 6 t
ORIGIN

Query Match 0.9%; Score 23; DB 6; Length 31;
Best Local Similarity 100.0%; Pred. No. 1.1;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1112 tcaacgccgaccggttcgtgca 1134
|||||
Db 23 TCAACGCCGACCGGTTCTTGCA 1

RESULT 10
AR092050 AR092050 31 bp DNA PAT 08-SEP-2000
LOCUS
DEFINITION Sequence 74 from patent US 5998141.
ACCESSION AR092050
VERSION AR092050.1 GI:10018804
KEYWORDS
SOURCE Unknown.
ORGANISM
REFERENCE Unclassified.
AUTHORS 1 (bases 1 to 31)
TITLE Acton,S.Laurene.
Intronic and polymorphic SR-BI nucleic acids and uses therefor

JOURNAL Patent: US 5998141-A 74 07-DEC-1999;
FEATURES Location/Qualifiers
Source 1..31
/organism="unknown"

BASE COUNT 6 a 12 c 6 g 7 t
ORIGIN

Query Match 0.9%; Score 23; DB 6; Length 31;
Best Local Similarity 100.0%; Pred. No. 1.1;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1112 tcaacgccgaccggttcgtgca 1134
|||||
Db 9 TCAACGCCGACCGGTTCTTGCA 31

RESULT 11
AR112183/c. AR112183 31 bp DNA PAT 16-MAY-2001
LOCUS
DEFINITION Sequence 72 from patent US 6130041.
ACCESSION AR112183
VERSION AR112183.1 GI:14092083

KEYWORDS
SOURCE Unknown.
ORGANISM Unclassified.
REFERENCE 1 (bases 1 to 31)
AUTHORS Acton,S.Laurene.
TITLE Human intronic and polymorphic SR-BI nucleic acids and uses therefor
JOURNAL Patent: US 6130041-A 72 10-OCT-2000;
FEATURES Location/Qualifiers
Source 1..31
/organism="unknown"
BASE COUNT 7 a 6 c 12 g 6 t
ORIGIN

Query Match 0.9%; Score 23; DB 6; Length 31;
Best Local Similarity 100.0%; Pred. No. 1.1;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1112 tcaacgccgacccggtctcgca 1134
DB 23 TCAACGCCGACCCGCTCTGCA 1

RESULT 12
LOCUS AR112185 31 bp DNA PAT 16-MAY-2001
DEFINITION Sequence 74 from patent US 6130041.
ACCESSION AR112185
VERSION AR112185.1 GI:14092085
KEYWORDS
SOURCE Unknown.
ORGANISM Unclassified.
REFERENCE 1 (bases 1 to 31)
AUTHORS Acton,S.Laurene.
TITLE Human intronic and polymorphic SR-BI nucleic acids and uses therefor
JOURNAL Patent: US 6130041-A 74 10-OCT-2000;
FEATURES Location/Qualifiers
Source 1..31
/organism="unknown"
BASE COUNT 6 a 12 c 6 g 7 t
ORIGIN

Query Match 0.9%; Score 23; DB 6; Length 31;
Best Local Similarity 100.0%; Pred. No. 1.1;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1112 tcaacgccgacccggtctcgca 1134
DB 9 TCAACGCCGACCCGCTCTGCA 31

RESULT 13
LOCUS AR149225 31 bp DNA PAT 08-AUG-2001
DEFINITION Sequence 72 from patent US 6228581.
ACCESSION AR149225
VERSION AR149225.1 GI:15113816
KEYWORDS
SOURCE Unknown.
ORGANISM Unclassified.
REFERENCE 1 (bases 1 to 31)
AUTHORS Acton,S.L. and Ordovas,J.M.
TITLE Human intronic and polymorphic SR-BI nucleic acids and uses therefor
JOURNAL Patent: US 6228581-A 72 08-MAY-2001;
FEATURES Location/Qualifiers
Source 1..31

BASE COUNT 7 a 6 c 12 g 6 t
ORIGIN

Query Match 0.9%; Score 23; DB 6; Length 31;
Best Local Similarity 100.0%; Pred. No. 1.1;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1112 tcaacgccgacccggtctcgca 1134
DB 23 TCAACGCCGACCCGCTCTGCA 1

RESULT 14
LOCUS AR149227 31 bp DNA PAT 08-AUG-2001
DEFINITION Sequence 74 from patent US 6228581.
ACCESSION AR149227
VERSION AR149227.1 GI:15113818
KEYWORDS
SOURCE Unknown.
ORGANISM Unclassified.
REFERENCE 1 (bases 1 to 31)
AUTHORS Acton,S.L. and Ordovas,J.M.
TITLE Human intronic and polymorphic SR-BI nucleic acids and uses therefor
JOURNAL Patent: US 6228581-A 74 08-MAY-2001;
FEATURES Location/Qualifiers
Source 1..31
/organism="unknown"
BASE COUNT 6 a 12 c 6 g 7 t
ORIGIN

Query Match 0.9%; Score 23; DB 6; Length 31;
Best Local Similarity 100.0%; Pred. No. 1.1;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1112 tcaacgccgacccggtctcgca 1134
DB 9 TCAACGCCGACCCGCTCTGCA 31

RESULT 15
LOCUS AR096475 36 bp DNA PAT 08-SEP-2000
DEFINITION Sequence 4 from patent US 6008014.
ACCESSION AR096475
VERSION AR096475.1 GI:10025310
KEYWORDS
SOURCE Unknown.
ORGANISM Unclassified.
REFERENCE 1 (bases 1 to 36)
AUTHORS Gimeno,C.J. and Acton,S.
TITLE Method of making lipid metabolic pathway compositions
JOURNAL Patent: US 6008014-A 4 28-DEC-1999;
FEATURES Location/Qualifiers
Source 1..36
/organism="unknown"
BASE COUNT 10 a 9 c 10 g 7 t
ORIGIN

Query Match 0.7%; Score 18; DB 6; Length 36;
Best Local Similarity 100.0%; Pred. No. 6.5e+02;
Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1456 caatccgagcccaag 1473
DB 19 CAATCCGAGCCCAAG 36

RESULT 16
AX161232/C
LOCUS AX161232 50 bp DNA PAT 22-JUN-2001
DEFINITION Sequence 4560 from Patent WO0140521.
ACCESSION AX161232
VERSION AX161232.1 GI:14542563
KEYWORDS
SOURCE human.
ORGANISM Homo sapiens
REFERENCE Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Homidae; Homo.
AUTHORS 1 (bases 1 to 50)
TITLE Shinkets, R.A. and Leach, M.
Nucleic acids containing single nucleotide polymorphisms and
methods of use thereof
JOURNAL Patent: WO 0140521-A 4560 07-JUN-2001;
Curagen Corporation (US)
FEATURES
source 1..50 Location/Qualifiers
misc_feature /organism="Homo sapiens"
/db_xref="taxon:9606"
25..26
/note="Nucleotide deleted between bases 25 and 26"
Accession number c943958770"
misc_feature 26
/note="2 of 2 allelic variants (4559 is other entry)"
BASE COUNT 16 a 4 c 5 g 25 t
ORIGIN

Query Match 0.7%; Score 17; DB 6; Length 50;
Best Local Similarity 100.0%; Pred. No. 6.4e+02;
Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2544 aaaaaatggaataaaaaa 2561
|||||
Db - 43 AAAAAATGCAAAAAAAA 26

RESULT 17
AR092047/C
LOCUS AR092047 20 bp DNA PAT 08-SEP-2000
DEFINITION Sequence 71 from patent US 5998141.
ACCESSION AR092047
VERSION AR092047.1 GI:10018801
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE Unclassified.
1 (bases 1 to 20)
AUTHORS Acton, S. Laurene.
TITLE Intronc and polymorphic SR-BI nucleic acids and uses therefor
JOURNAL Patent: US 5998141-A 71 07-DEC-1999;
FEATURES Location/Qualifiers
source 1..20
/organism="unknown"
BASE COUNT 4 a 4 c 8 g 4 t
ORIGIN

Query Match 0.7%; Score 17; DB 6; Length 20;
Best Local Similarity 100.0%; Pred. No. 2.3e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgcgcacccggtt 1128
|||||
Db 17 TCAACGCCGACCCGGTT 1

RESULT 18
AR092049

LOCUS AR092049 20 bp DNA PAT 08-SEP-2000
DEFINITION Sequence 73 from patent US 5998141.
ACCESSION AR092049
VERSION AR092049.1 GI:10018803
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE Unclassified.
1 (bases 1 to 20)
AUTHORS Acton, S. Laurene.
TITLE Intronc and polymorphic SR-BI nucleic acids and uses therefor
JOURNAL Patent: US 5998141-A 73 07-DEC-1999;
FEATURES Location/Qualifiers
source 1..20
/organism="unknown"
BASE COUNT 4 a 8 c 4 g 4 t
ORIGIN

Query Match 0.7%; Score 17; DB 6; Length 20;
Best Local Similarity 100.0%; Pred. No. 2.3e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgcgcacccggtt 1128
|||||
Db 4 TCAACGCCGACCCGGTT 20

RESULT 19
AR112182/C
LOCUS AR112182 20 bp DNA PAT 16-MAY-2001
DEFINITION Sequence 71 from patent US 6130041.
ACCESSION AR112182
VERSION AR112182.1 GI:14092082
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE Unclassified.
1 (bases 1 to 20)
AUTHORS Acton, S. Laurene.
TITLE Human Intronc and polymorphic SR-BI nucleic acids and uses therefor
JOURNAL Patent: US 6130041-A 71 10-OCT-2000;
FEATURES Location/Qualifiers
source 1..20
/organism="unknown"
BASE COUNT 4 a 4 c 8 g 4 t
ORIGIN

Query Match 0.7%; Score 17; DB 6; Length 20;
Best Local Similarity 100.0%; Pred. No. 2.3e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgcgcacccggtt 1128
|||||
Db 17 TCAACGCCGACCCGGTT 1

RESULT 20
AR112184
LOCUS AR112184 20 bp DNA PAT 16-MAY-2001
DEFINITION Sequence 73 from patent US 6130041.
ACCESSION AR112184
VERSION AR112184.1 GI:14092084
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE Unclassified.
1 (bases 1 to 20)
AUTHORS Acton, S. Laurene.
TITLE Human Intronc and polymorphic SR-BI nucleic acids and uses therefor

JOURNAL Patent: US 6130041-A 73 10-OCT-2000;
FEATURES Location/Qualifiers
source 1..20
/organism="unknown"
BASE COUNT 4 a 8 c 4 g 4 t
ORIGIN

Query Match 0.7%; Score 17; DB 6; Length 20;
Best Local Similarity 100.0%; Pred. No. 2.3e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgccgacccggtt 1128
|||||
Db 4 TCAACGCCGACCCGTT 20

RESULT 21
LOCUS ARI49224 20 bp DNA PAT 08-AUG-2001
DEFINITION Sequence 71 from patent US 6228581.
ACCESSION ARI49224
VERSION ARI49224.1 GI:15113815
KEYWORDS
SOURCE Unknown.
ORGANISM Unclassified.

REFERENCE 1 (bases 1 to 20)
AUTHORS Acton,S.L. and Ordovas,J.M.
TITLE Human intronic and polymorphic SR-BI nucleic acids and uses.
JOURNAL Patent: US 6228581-A 71 08-MAY-2001;
FEATURES Location/Qualifiers
source 1..20
/organism="unknown"

BASE COUNT 4 a 4 c 8 g 4 t
ORIGIN

Query Match 0.7%; Score 17; DB 6; Length 20;
Best Local Similarity 100.0%; Pred. No. 2.3e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgccgacccggtt 1128
|||||
Db 17 TCAACGCCGACCCGTT 1

RESULT 22
LOCUS ARI49226 20 bp DNA PAT 08-AUG-2001
DEFINITION Sequence 73 from patent US 6228581.
ACCESSION ARI49226
VERSION ARI49226.1 GI:15113817
KEYWORDS
SOURCE Unknown.
ORGANISM Unclassified.

REFERENCE 1 (bases 1 to 20)
AUTHORS Acton,S.L. and Ordovas,J.M.
TITLE Human intronic and polymorphic SR-BI nucleic acids and uses.
JOURNAL Patent: US 6228581-A 73 08-MAY-2001;
FEATURES Location/Qualifiers
source 1..20
/organism="unknown"

BASE COUNT 4 a 8 c 4 g 4 t
ORIGIN

Query Match 0.7%; Score 17; DB 6; Length 20;
Best Local Similarity 100.0%; Pred. No. 2.3e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1112 tcaacgccgacccggtt 1128
|||||
Db 4 TCAACGCCGACCCGTT 20

RESULT 23
LOCUS AX043026/c 25 bp DNA PAT 23-NOV-2000
DEFINITION Sequence 592 from Patent WO0065088.
ACCESSION AX043026
VERSION AX043026.1 GI:11341634
KEYWORDS
SOURCE synthetic construct.
ORGANISM synthetic construct.
REFERENCE 1 (bases 1 to 25)
AUTHORS Ulfendahl,P.J. and Wong,K.C.
TITLE Primers for identifying typing or classifying nucleic acids
JOURNAL Patent: WO 0065088-A 592 02-NOV-2000;
FEATURES Location/Qualifiers
source 1..25
/organism="synthetic construct"
/db_xref="taxon:32630"
/note="16S rRNA Homozygote Primer Sequence"

BASE COUNT 2 a 3 c 3 g 17 t
ORIGIN

Query Match 0.7%; Score 17; DB 6; Length 25;
Best Local Similarity 100.0%; Pred. No. 2.3e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2549 atgggaaaaaataaaa 2565
|||||
Db 17 ATGGGAAAAAATAAAA 1

RESULT 24
LOCUS ARI42905 47 bp DNA PAT 08-AUG-2001
DEFINITION Sequence 1 from patent US 6204024.
ACCESSION ARI42905
VERSION ARI42905.1 GI:15104191
KEYWORDS
SOURCE Unknown.
ORGANISM Unclassified.

REFERENCE 1 (bases 1 to 47)
AUTHORS Romano,J.W. and Lee,E.M.
TITLE CCR5 RNA transcription based amplification assay
JOURNAL Patent: US 6204024-A 1 20-MAR-2001;
FEATURES Location/Qualifiers
source 1..47
/organism="unknown"

BASE COUNT 15 a 14 c 11 g 7 t
ORIGIN

Query Match 0.7%; Score 17; DB 6; Length 47;
Best Local Similarity 100.0%; Pred. No. 2.3e+03;
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1295 tggctccgacgctgctc 1311
|||||
Db 41 TGGTCTCGACCGCTCTC 25

RESULT 25
LOCUS E32461 18 bp DNA PAT 07-FEB-2001
DEFINITION Mammal-derived tissue specific physiologically active protein.

ACCESSION E32461
 VERSION E32461.1 GI:13018697
 KEYWORDS JP 2000037190-A/21.
 SOURCE unidentified.
 ORGANISM unidentified.
 REFERENCE 1 (bases 1 to 18)
 AUTHORS Jun,N.Y,N.N. and Tanaka.
 TITLE Mammal-derived tissue specific physiologically active protein
 JOURNAL Patent: JP 2000037190-A 21 08-FEB-2000;
 JAPAN TOBACCO INC
 COMMENT
 OS Artificial Sequence
 PN JP 2000037190-A/21
 PD 08-FEB-2000
 PE 23-JUL-1998 JP 1998225228
 PR JUN NISHIU,YOSUKE NAKAMURA,TOSHIHIRO TANAKA
 PC C12N15/09,C07K14/47,C07K16/18,C12N1/19,C12N1/21,C12N5/10, PC
 C12N15/02,
 PC C12P21/02,C12P21/08/(C12N5/10,C12R1:91),(C12P21/08,C12R1:91),
 PC C12N15/00,
 PC C12N5/00,C12N15/00,(C12N5/00,C12R1:91)
 CC
 C12N15/00,C12N15/00,(C12N5/00,C12R1:91)
 FT primer_bind
 Location/Qualifiers
 1..18
 /organism="unidentified"
 /db_xref="taxon:32644"
 BASE COUNT 0 a 2 c 1 g 15 t
 ORIGIN

Query Match 0.6%; Score 16; DB 6; Length 18;
 Best Local Similarity 100.0%; Pred. No. 8.4e+03;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 Oy 2551 ggaataaaaaa 2566
 |||||||
 Db 18 GGAATAAAAAA 3

RESULT 26
 E29883/c
 LOCUS E29883 20 bp DNA PAT 07-FEB-2001
 DEFINITION HIV cofactor inhibitor.
 E29883
 ACCESSION E29883.1 GI:13021278
 VERSION JP 1999292795-A/37.
 KEYWORDS
 SOURCE unidentified.
 ORGANISM unidentified.
 REFERENCE 1 (bases 1 to 20)
 AUTHORS Hiroshi,T.N.Y.Y. and Kimura,K.T.A.A.
 TITLE HIV cofactor inhibitor
 JOURNAL Patent: JP 1999292795-A 37 26-OCT-1999;
 YAMANOUCHI PHARMACEUT CO LTD
 COMMENT
 OS Unidentified
 PN JP 1999292795-A/37
 PD 26-OCT-1999
 PE 02-APR-1998 JP 1998125452
 PR HIROSHI TAKAHISA,NAOKI YAMAMOTO,TORU KIMURA,KAZUYUKI TAKAI, PI
 PC AKIRA WADA
 PC A61K48/00,A61K31/70,A61K31/70,C12N15/09,C12N15/00 CC
 FH Key Location/Qualifiers
 FT source 1..20
 Location/Qualifiers
 1..20
 /organism="unidentified"
 /db_xref="taxon:32644"
 BASE COUNT 5 a 8 c 7 g 0 t

ORIGIN
 Query Match 0.6%; Score 16; DB 6; Length 20;
 Best Local Similarity 100.0%; Pred. No. 8.3e+03;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 Oy 1295 tggctcgcgcgtct 1310
 |||||||
 Db 16 TGGCTCCTCCGCTCT 1

RESULT 27
 ARI42908/c
 LOCUS ARI42908 22 bp DNA PAT 08-AUG-2001
 DEFINITION Sequence 4 from patent US 6204024.
 ARI42908
 ACCESSION ARI42908.1 GI:15104194
 VERSION
 KEYWORDS
 SOURCE Unknown.
 ORGANISM Unknown.
 REFERENCE 1 (bases 1 to 22)
 AUTHORS Romano,J.W. and Lee,E.M.
 TITLE CCR5 RNA transcription based amplification assay
 JOURNAL Patent: US 6204024-A 4 20-MAR-2001;
 Location/Qualifiers
 1..22
 /organism="unknown"
 BASE COUNT 6 a 9 c 7 g 0 t
 ORIGIN

Query Match 0.6%; Score 16; DB 6; Length 22;
 Best Local Similarity 100.0%; Pred. No. 8.3e+03;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 Oy 1295 tggctcgcgcgtct 1310
 |||||||
 Db 16 TGGCTCCTCCGCTCT 1

RESULT 28
 AX115478
 LOCUS AX115478 23 bp DNA PAT 11-MAY-2001
 DEFINITION Sequence 601 from Patent WO0129262.
 AX115478
 ACCESSION AX115478.1 GI:14032420
 VERSION
 KEYWORDS
 SOURCE synthetic construct.
 ORGANISM synthetic construct.
 REFERENCE 1 (bases 1 to 23)
 AUTHORS Picoult-Newburg,L. and Pohl,M.
 TITLE Genotyping reagents, kits and methods of use thereof
 JOURNAL Patent: WO 0129262-A 601 26-APR-2001;
 Orchid Biosciences, Inc. (US)
 COMMENT
 OS Unidentified
 PN 1..23
 Location/Qualifiers
 1..23
 /organism="synthetic construct"
 /db_xref="taxon:32630"
 /note="Primer"
 BASE COUNT 16 a 1 c 5 g 1 t
 ORIGIN
 Query Match 0.6%; Score 16; DB 6; Length 23;
 Best Local Similarity 100.0%; Pred. No. 8.3e+03;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 Oy 2551 ggaataaaaaa 2566
 |||||||
 Db 5 GGAATAAAAAA 20

RESULT 29
LOCUS A90999 25 bp DNA PAT 22-JAN-2000
DEFINITION Sequence 1 from Patent EP0854196.
ACCESSION A90999
VERSION A90999.1 GI:6739605
KEYWORDS
SOURCE unidentified.
ORGANISM unidentified.
REFERENCE 1 (bases 1 to 25)
AUTHORS Paabeo,S.E. and Kilger,C.A.
TITLE Method for the uncoupled, direct, exponential amplification and sequencing of DNA molecules with the addition of a second thermostable DNA polymerase and its application
JOURNAL Patent: EP 0854196-A 1 22-JUL-1998;
BOEHRINGER MANNHEIM GMBH (DE)
FEATURES
source Location/Qualifiers
1..25
/organism="unidentified"
/db_xref="taxon:32644"
BASE COUNT 1 a 8 c 8 g 8 t
ORIGIN

Query Match 0.6%; Score 16; DB 6; Length 25;
Best Local Similarity 100.0%; Pred. No. 8.3e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1295 tggctcgcgcgcgtcgt 1310
|||||
Db 4 TGGTCTGCGCGCTGCT 19

RESULT 30
LOCUS A91901 25 bp DNA PAT 22-JAN-2000
DEFINITION Sequence 5 from Patent EP0849364.
ACCESSION A91901
VERSION A91901.1 GI:6740774
KEYWORDS
SOURCE unidentified.
ORGANISM unidentified.
REFERENCE 1 (bases 1 to 25)
AUTHORS Paabeo,S.E. and Kilger,C.A.
TITLE Method for the direct, exponential amplification and sequencing of DNA molecules and its application
JOURNAL Patent: EP 0849364-A 5 24-JUN-1998;
BOEHRINGER MANNHEIM GMBH (DE)
FEATURES
source Location/Qualifiers
1..25
/organism="unidentified"
/db_xref="taxon:32644"
BASE COUNT 1 a 8 c 8 g 8 t
ORIGIN

Query Match 0.6%; Score 16; DB 6; Length 25;
Best Local Similarity 100.0%; Pred. No. 8.3e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1295 tggctcgcgcgcgtcgt 1310
|||||
Db 4 TGGTCTGCGCGCTGCT 19

RESULT 31
LOCUS ARI06367 25 bp DNA PAT 14-FEB-2001
DEFINITION Sequence 5 from patent US-6107032.

ACCESSION ARI06367
VERSION ARI06367.1 GI:12820897
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE 1 (bases 1 to 25)
AUTHORS Kilger,C. and Paabo,S.
TITLE Method for the direct, exponential amplification and sequencing of DNA molecules and its application
JOURNAL Patent: US 6107032-A 5 22-AUG-2000;
FEATURES
source Location/Qualifiers
1..25
/organism="unknown"
BASE COUNT 1 a 8 c 8 g 8 t
ORIGIN

Query Match 0.6%; Score 16; DB 6; Length 25;
Best Local Similarity 100.0%; Pred. No. 8.3e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1295 tggctcgcgcgcgtcgt 1310
|||||
Db 4 TGGTCTGCGCGCTGCT 19

RESULT 32
LOCUS ARI48371 25 bp DNA PAT 08-AUG-2001
DEFINITION Sequence 1 from patent US 6225092.
ACCESSION ARI48371
VERSION ARI48371.1 GI:15112461
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE 1 (bases 1 to 25)
AUTHORS Kilger,C. and Paabo,S.
TITLE Method for the uncoupled, direct, exponential amplification and sequencing of DNA molecules with the addition of a second thermostable DNA polymerase and its application
JOURNAL Patent: US 6225092-A 1 01-MAY-2001;
FEATURES
source Location/Qualifiers
1..25
/organism="unknown"
BASE COUNT 1 a 8 c 8 g 8 t
ORIGIN

Query Match 0.6%; Score 16; DB 6; Length 25;
Best Local Similarity 100.0%; Pred. No. 8.3e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1295 tggctcgcgcgcgtcgt 1310
|||||
Db 4 TGGTCTGCGCGCTGCT 19

RESULT 33
LOCUS AX032404 25 bp DNA PAT 20-SEP-2000
DEFINITION Sequence 5 from Patent EP1004677.
ACCESSION AX032404
VERSION AX032404.1 GI:10279377
KEYWORDS
SOURCE unidentified.
ORGANISM unidentified.
REFERENCE 1 (bases 1 to 25)
AUTHORS Paabeo,S.E. and Kilger,C.A.
TITLE Method for the direct, exponential amplification and

JOURNAL Patent: EP 1004677-A 5 31-MAY-2000;

ROCHE DIAGNOSTICS GMBH (DE)
FEATURES

source

1.25
/organism="unidentified"

/db_xref="taxon:32630"

BASE COUNT 1 a 8 c 8 g 8 t

ORIGIN

Query Match

0.6%; Score 16; DB 6; Length 25;

Best Local Similarity 100.0%; Pred. No. 8.3e+03; Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1295 tggctcctgcgcctgct 1310

DB 4 TGCTCTCGCCCTGCT 19

RESULT 34

AX042532/C

25 bp DNA

PAT 23-NOV-2000

DEFINITION Sequence 98 from Patent WO0065088.

ACCESSION AX042532

GI:11341140

VERSION

KEYWORDS

SOURCE

ORGANISM

REFERENCE

AUTHORS

TITLE

JOURNAL

Amersham Pharmacia Biotech AB (SE)

Location/Qualifiers

FEATURES

source

1.25

/organism="synthetic construct"

/db_xref="taxon:32630"

/note="DPBI Homozygote primer sequence"

BASE COUNT 2 a 6 c 1 g

ORIGIN

Query Match

Best Local Similarity 100.0%; Score 16; DB 6; Length 25;

Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2550 tggaaaaa

DB 16 TCGAAAAA

RESULT 35

AX042542/C

25 bp DNA

PAT 23-NOV-2000

DEFINITION Sequence 108 from Patent WO0065088.

ACCESSION AX042542

VERSION AX042542.1

KEYWORDS

SOURCE

ORGANISM

REFERENCE

AUTHORS

TITLE

JOURNAL

Amersham Pharmacia Biotech AB (SE)

Location/Qualifiers

FEATURES

source

1.25

/organism="synthetic construct"

/db_xref="taxon:32630"

/note="DPBI Homozygote primer sequence"

BASE COUNT 5 a 3 c 2 g 15 t

ORIGIN

Query Match 0.6%; Score 16; DB 6; Length 25;
Best Local Similarity 100.0%; Pred. No. 8.3e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2550 tggaaaaa

DB 16 TCGAAAAA

RESULT 36

AX042571/C

25 bp DNA

PAT 23-NOV-2000

DEFINITION Sequence 137 from Patent WO0065088.

ACCESSION AX042571

VERSION AX042571.1

KEYWORDS

SOURCE

ORGANISM

REFERENCE

AUTHORS

TITLE

JOURNAL

Amersham Pharmacia Biotech AB (SE)

Location/Qualifiers

FEATURES

source

1.25

/organism="synthetic construct"

/db_xref="taxon:32630"

/note="DPBI Homozygote primer sequence"

BASE COUNT 2 a 6 c 2 g 15 t

ORIGIN

Query Match

Best Local Similarity 100.0%; Score 16; DB 6; Length 25;

Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2550 tggaaaaa

DB 16 TCGAAAAA

RESULT 37

AX042589/C

25 bp DNA

PAT 23-NOV-2000

DEFINITION Sequence 155 from Patent WO0065088.

ACCESSION AX042589

VERSION AX042589.1

KEYWORDS

SOURCE

ORGANISM

REFERENCE

AUTHORS

TITLE

JOURNAL

Amersham Pharmacia Biotech AB (SE)

Location/Qualifiers

FEATURES

source

1.25

/organism="synthetic construct"

/db_xref="taxon:32630"

/note="DPBI Homozygote primer sequence"

BASE COUNT 0 a 6 c 1 g 18 t

ORIGIN

Query Match 0.6%; Score 16; DB 6; Length 25;
Best Local Similarity 100.0%; Pred. No. 8.3e+03;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2551 gga

Db 16 GGAAGAAAAA 1

RESULT 38
LOCUS AX042627/c 25 bp DNA PAT 23-NOV-2000
DEFINITION Sequence 193 from Patent: WO0065088.
ACCESSION AX042627
VERSION AX042627.1 GI:11341235
KEYWORDS
SOURCE synthetic construct.
ORGANISM synthetic construct.
REFERENCE 1 (bases 1 to 25)
AUTHORS Ulfendahl, P.J. and Wong, K.C.
TITLE Primers for identifying typing or classifying nucleic acids
JOURNAL Patent: WO 0065088-A 193 02-NOV-2000;
Amersham Pharmacia Biotech AB (SE)
FEATURES
source 1..25
/organism="synthetic construct"
/db_xref="taxon:32630"
/note="HLA-B Homozygote Primer Sequence"

BASE COUNT 2 a 5 c 4 g 14 t

ORIGIN

Query Match
Best Local Similarity 100.0%; Score 16; DB 6; Length 25;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2550 tggagaaaaa 2565
Db 16 TGGAAAAA 1

RESULT 39
LOCUS AX042759/c 25 bp DNA PAT 23-NOV-2000
DEFINITION Sequence 325 from Patent: WO0065088.
ACCESSION AX042759
VERSION AX042759.1 GI:11341367
KEYWORDS
SOURCE synthetic construct.
ORGANISM synthetic construct.
REFERENCE 1 (bases 1 to 25)
AUTHORS Ulfendahl, P.J. and Wong, K.C.
TITLE Primers for identifying typing or classifying nucleic acids
JOURNAL Patent: WO 0065088-A 325 02-NOV-2000;
Amersham Pharmacia Biotech AB (SE)
FEATURES
source 1..25
/organism="synthetic construct"
/db_xref="taxon:32630"
/note="HLA-B Homozygote Primer Sequence"

BASE COUNT 3 a 2 c 4 g 16 t

ORIGIN

Query Match
Best Local Similarity 100.0%; Score 16; DB 6; Length 25;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2550 tggagaaaaa 2565
Db 16 TGGAAAAA 1

RESULT 40
LOCUS AX042823/c 25 bp DNA PAT 23-NOV-2000

DEFINITION Sequence 389 from Patent: WO0065088.
ACCESSION AX042823
VERSION AX042823.1 GI:11341431
KEYWORDS
SOURCE synthetic construct.
ORGANISM synthetic construct.
REFERENCE 1 (bases 1 to 25)
AUTHORS Ulfendahl, P.J. and Wong, K.C.
TITLE Primers for identifying typing or classifying nucleic acids
JOURNAL Patent: WO 0065088-A 389 02-NOV-2000;
Amersham Pharmacia Biotech AB (SE)
FEATURES
source 1..25
/organism="synthetic construct"
/db_xref="taxon:32630"
/note="HLA-B Homozygote Primer Sequence"

BASE COUNT 3 a 7 c 0 g 15 t

ORIGIN

Query Match
Best Local Similarity 100.0%; Score 16; DB 6; Length 25;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaagaaaaa 2566
Db 16 GGAAGAAAAA 1

RESULT 41
LOCUS AX042825/c 25 bp DNA PAT 23-NOV-2000
DEFINITION Sequence 391 from Patent: WO0065088.
ACCESSION AX042825
VERSION AX042825.1 GI:11341433
KEYWORDS
SOURCE synthetic construct.
ORGANISM synthetic construct.
REFERENCE 1 (bases 1 to 25)
AUTHORS Ulfendahl, P.J. and Wong, K.C.
TITLE Primers for identifying typing or classifying nucleic acids
JOURNAL Patent: WO 0065088-A 391 02-NOV-2000;
Amersham Pharmacia Biotech AB (SE)
FEATURES
source 1..25
/organism="synthetic construct"
/db_xref="taxon:32630"
/note="HLA-B Homozygote Primer Sequence"

BASE COUNT 2 a 7 c 1 g 15 t

ORIGIN

Query Match
Best Local Similarity 100.0%; Score 16; DB 6; Length 25;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaagaaaaa 2566
Db 16 GGAAGAAAAA 1

RESULT 42
LOCUS AX042827/c 25 bp DNA PAT 23-NOV-2000
DEFINITION Sequence 393 from Patent: WO0065088.
ACCESSION AX042827
VERSION AX042827.1 GI:11341435
KEYWORDS
SOURCE synthetic construct.
ORGANISM synthetic construct.
REFERENCE 1 (bases 1 to 25)
AUTHORS Ulfendahl, P.J. and Wong, K.C.
TITLE Primers for identifying typing or classifying nucleic acids
JOURNAL Patent: WO 0065088-A 393 02-NOV-2000;
Amersham Pharmacia Biotech AB (SE)
FEATURES
source 1..25
/organism="synthetic construct"
/db_xref="taxon:32630"
/note="HLA-B Homozygote Primer Sequence"

BASE COUNT 2 a 7 c 1 g 15 t

ORIGIN

Query Match
Best Local Similarity 100.0%; Score 16; DB 6; Length 25;
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2551 ggaagaaaaa 2566
Db 16 GGAAGAAAAA 1

REFERENCE 1 (bases 1 to 25)
 AUTHORS Ulfendahl, P.J. and Wong, K.C.
 TITLE Primers for identifying typing or classifying nucleic acids
 JOURNAL Patent: WO 0055088-A 393 02-NOV-2000;
 Amersham Pharmacia Biotech AB (SE)

FEATURES
 source 1..25
 /organism="synthetic construct"
 /db_xref="taxon:32630"
 /note="HLA-B Homozygote Primer Sequence"

BASE COUNT 2 a 7 c 2 g 14 t
 ORIGIN

Query Match 0.6%; Score 16; DB 6; Length 25;
 Best Local Similarity 100.0%; Pred. No. 8.3e+03;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 2550 tggaaaaa 2565
 |||||
 Db 16 TGGAAAAA 1

RESULT 43
 LOCUS AX042871 25 bp DNA PAT 23-NOV-2000
 DEFINITION Sequence 437 from Patent WO0065088.
 ACCESSION AX042871
 VERSION AX042871.1 GI:11341479.
 KEYWORDS
 SOURCE synthetic construct.
 ORGANISM artificial sequence.

REFERENCE 1 (bases 1 to 25)
 AUTHORS Ulfendahl, P.J. and Wong, K.C.
 TITLE Primers for identifying typing or classifying nucleic acids
 JOURNAL Patent: WO 0065088-A 437 02-NOV-2000;
 Amersham Pharmacia Biotech AB (SE)

FEATURES
 source 1..25
 /organism="synthetic construct"
 /db_xref="taxon:32630"
 /note="HLA-C Homozygote Primer Sequence"

BASE COUNT 5 a 2 c 2 g 16 t
 ORIGIN

Query Match 0.6%; Score 16; DB 6; Length 25;
 Best Local Similarity 100.0%; Pred. No. 8.3e+03;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 2550 tggaaaaa 2565
 |||||
 Db 16 TGGAAAAA 1

RESULT 44
 LOCUS AX042878 25 bp DNA PAT 23-NOV-2000
 DEFINITION Sequence 444 from Patent WO0065088.
 ACCESSION AX042878
 VERSION AX042878.1 GI:11341486
 KEYWORDS
 SOURCE synthetic construct.
 ORGANISM artificial sequence.

REFERENCE 1 (bases 1 to 25)
 AUTHORS Ulfendahl, P.J. and Wong, K.C.
 TITLE Primers for identifying typing or classifying nucleic acids
 JOURNAL Patent: WO 0065088-A 444 02-NOV-2000;
 Amersham Pharmacia Biotech AB (SE)

FEATURES
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 Location/Qualifiers

/organism="synthetic construct"
 /db_xref="taxon:32630"
 /note="HLA-C Homozygote Primer Sequence"

BASE COUNT 3 a 2 c 4 g 16 t
 ORIGIN

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 Best Local Similarity 100.0%; Pred. No. 8.3e+03;
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

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 Db 16 TGGAAAAA 1

RESULT 45
 LOCUS AX032410 25 bp DNA UNA 20-SEP-2000
 DEFINITION Sequence 5 from Patent EP1004677.
 ACCESSION AX032410
 VERSION AX032410.1 GI:10279383
 KEYWORDS
 SOURCE unidentified.
 ORGANISM unidentified.

REFERENCE 1 (bases 1 to 25)
 AUTHORS Paeaebo, S.E. and Kilger, C.A.
 TITLE Method for the direct, exponential amplification and
 JOURNAL Patent: EP 1004677-A 31-MAY-2000;
 ROCHE DIAGNOSTICS GMBH (DE)

FEATURES
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 /organism="unidentified"
 /db_xref="taxon:32644"
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 ORIGIN

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 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 1295 tggctcgtgcgcgtgct 1310
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 Db 4 TGGTCTGCGCGTCT 19

Search completed: April 20, 2002, 10:07:59
 Job time: 12206 sec

